

ORF predicted by Cole et al. (Nature 393:537-544) and containing Rv0546c

1/1	31/11
tag tca ggg cgt gca ttc gac gac gct gta	cta ccc gct ggt ggc aac tcc gat gat tgc
AMB ser gly arg ala phe asp asp ala val	leu pro ala gly gly asn ser asp asp cys
61/21	91/31
	get gac ege ege gga age ege ega gtg gat
	ala asp arg arg gly ser arg arg val asp
121/41	151/51
	gat cgc gcc tcg ggt tgc cgt cgc cgt caa asp arg ala ser gly cys arg arg arg gln
181/61	211/71
	caa tgc act cat gca gcg ccg caa cga aca
	gln cys thr his ala ala pro gln arg thr
241/81	271/91
gct caa ccc ttg aac cgg gtc ccg gcc tgc	cga ccc tcg gcc gcc ggc gtg ccg cta cgt
ala gln pro leu asn arg val pro ala cys	arg pro ser ala ala gly val pro leu arg
301/101	331/111
	age egg atg eta ett egg eeg geg gae tat
	ser arg met leu leu arg pro ala asp tyr
361/121	391/131
	atc ggg ctg gcg att gcc cgt gaa tac ggg
	ile gly leu ala ile ala arg glu tyr gly 451/151
421/141	ctg ctc gaa ctg gcc ggt tac ggc gag ccg
	leu leu glu leu ala gly tyr gly glu pro
481/161	511/171
	ctg tgg ctg cag gtg cgc gac ctc gag gct
	leu trp leu gln val arg asp leu glu ala
541/181	571/191
acc cag acc gag ctg gtc agc cga ggc gtg	tcg atc gct cgc gag ccc cgc cgc gaa ccg
	ser ile ala arg glu pro arg arg glu pro
601/201	631/211
	cca gac ggg atc aca ctg ata ttc gtc gag
	pro asp gly ile thr leu ile phe val glu
661/221	691/231
gtt ccc gag ggt cac ccg ctg cgt aca gac	
val pro glu gly his pro leu arg thr asp	thr arg ala OPA

SEQ ID No.11F

FIGURE 11F

31/11 1/1 qac cga agg gat ttc gcg act aac tcg gcc tgt aag gca acg cga ggt ctt cat gcc gag asp arg arg asp phe ala thr asn ser ala cys lys ala thr arg gly leu his ala glu 91/31 gac gta gac agg aag aga cag gga agc tga tga cgt cgc gta ccg gac cgc cat tct gtc asp val asp arg lys arg gln gly ser OPA OPA arg arg val pro asp arg his ser val 151/51 121/41 gag tot tto oga gtt cag caa caa tog aca cag aag ogg gga oca gac ogg gag gac gac glu ser phe arg val gln gln ser thr gln lys arg gly pro asp arg glu asp asp 211/71 geg gee egg gee get teg gge ega gtg tet gag taa gae eag agt eae ggg tee gtg tgt ala ala arg ala ala ser gly arg val ser glu OCH asp gln ser his gly ser val cys 271/91 qac aac cgc gcg gaa ttc aat cgg atg gcg ggc ggg acc gga ttg cgc cgg tca ccg agg asp asn arg ala glu phe asn arg met ala gly gly thr gly leu arg arg ser pro arg aac ctc cgg agt gat c asn leu arg ser asp

SEQ ID No.12A

FIGURE 12A

31/11 acc gaa ggg att tcg cga cta act cgg cct gta agg caa cgc gag gtc ttc atg ccg agg thr glu gly ile ser arg leu thr arg pro val arg gln arg glu val phe met pro arg 91/31 acg tag aca gga aga gac agg gaa gct gat gac gtc gcg tac cgg acc gcc att ctg tcg thr AMB thr gly arg asp arg glu ala asp asp val ala tyr arg thr ala ile leu ser 151/51 ser leu ser glu phe ser asn asn arg his arg ser gly asp gln thr gly arg thr thr 211/71 cgg ccc ggg ccg ctt cgg gcc gag tgt ctg agt aag acc aga gtc acg ggt ccg tgt gtg arg pro gly pro leu arg ala glu cys leu ser lys thr arg val thr gly pro cys val 271/91 241/81 aca acc gcg cgg aat tca atc gga tgg cgg gcg gga ccg gat tgc gcc ggt cac cga gga thr thr ala arg asn ser ile gly trp arg ala gly pro asp cys ala gly his arg gly 301/101 acc tcc gga gtg atc thr ser gly val ile

SEQ ID No.12B

FIGURE 12B

ccg aag gga ttt cgc gac taa ctc ggc ctg taa ggc aac gcg agg tct tca tgc cga gga pro lys gly phe arg asp OCH leu gly leu OCH gly asn ala arg ser ser cys arg gly 91/31 61/21 cqt aga cag gaa gag aca ggg aag ctg atg acg tcg cgt acc gga ccg cca ttc tgt cga arg arg gln glu glu thr gly lys leu met thr ser arg thr gly pro pro phe cys arg 121/41 val phe pro ser ser ala thr ile asp thr glu ala gly thr arg pro gly gly arg arg 211/71 181/61 ggc ccg ggc cgc ttc ggg ccg agt gtc tga gta aga cca gag tca cgg gtc cgt gtg tga gly pro gly arg phe gly pro ser val OPA val arg pro glu ser arg val arg val OPA 271/91 241/81 caa ccg cgc gga att caa tcg gat ggc ggg cgg gac cgg att gcg ccg gtc acc gag gaa gln pro arg gly ile gln ser asp gly gly arg asp arg ile ala pro val thr glu glu 301/101 cct ccg gag tga tc pro pro glu OPA

SEO ID No.12C

FIGURE 12C

31/11 1/1 GGG ATT TCG TTG CCC GAT GGA TTG TTT GTA CGG TTT GGG AAA AAC ACT TGA AGT CCT TTT gly ile ser leu pro asp gly leu phe val arg phe gly lys asn thr OPA ser pro phe 61/21 91/31 TAT TGG CAA TGC TGG AAA TGG ACA TTC CAA TAT TGC GCG AAT TAA CCG AAC ACG GTG AGG tyr trp gln cys trp lys trp thr phe gln tyr cys ala asn OCH pro asn thr val arg 151/51 121/41 GGG GGG CAA GCG TTT GTA CCG GGG CCA GCA AGC GCC GCC GAC CGG TTG ACC GAA GCC AGC gly gly gln ala phe val pro gly pro ala ser ala ala asp arg leu thr glu ala ser 211/71 ATG TTG TTG TGT CAG CGC GGG CTT GGT CTC GAT GTC CCG GCC TTG GCT GGA CCC GCT TCT met leu cys gln arg gly leu gly leu asp val pro ala leu ala gly pro ala ser 271/91 241/81 TCA AAA CAG GTT GAA CTT AAC GAC TCA AGA ACG GAA ACG CTT GAA CCG CGA CGT CGC TCC ser lys gln val glu leu asn asp ser arg thr glu thr leu glu pro arg arg ser 301/101 331/111 GGA CAC CAA TTT GAC TCG GCT CTT TGG CAA TTG AAG GTG AGC TGC GAG CAG CCG GGT GAC gly his gln phe asp ser ala leu trp gln leu lys val ser cys glu gln pro gly asp 391/131 361/121 CGC ATC GTT GGC CTT GCC ATC AAT CGC CGG CTC GCG GAC GTA GAT AAT CAG CTC ACC GTT arg ile val gly leu ala ile asn arg arg leu ala asp val asp asn gln leu thr val 451/151 421/141 GGG ACC GAC CTC GAC CAG GGG TCC TTT GTG ACT GCC GGG CTT GAC GCG GAC GAC CAC AGA gly thr asp leu asp gln gly ser phe val thr ala gly leu asp ala asp asp his arg 511/171 481/161 GTC GGT CAT CGC CTA AGG CTA CCG TTC TGA CCT GGG GCT GCG TGG GCG CCG ACG ACG TGA val gly his arg leu arg leu pro phe OPA pro gly ala ala trp ala pro thr thr OPA 571/191 541/181 GGC ACG TCA TGT CTC AGC GGC CCA CCG CCA CCT CGG TCG CCG GCA GTA TGT CAG CAT GTG gly thr ser cys leu ser gly pro pro pro pro arg ser pro ala val cys gln his val 631/211 601/201 CAG ATG ACT CCA CGC AGC CTT GTT CGC ATC GTT GGT GTC GTG GTT GCG ACG ACC TTG GCG gln met thr pro arg ser leu val arg ile val gly val val val ala thr thr leu ala 691/231 661/221 CTG GTG AGC GCA CCC GCC GGC GGT CGT GCC GCG CAT GCG GAT C leu val ser ala pro ala gly gly arg ala ala his ala asp

SEQ ID No.13A

FIGURE 13A



	32/11
GGA TTT CGT TGC CCG ATG GAT TGT TTG TAC	GGT TTG GGA AAA ACA CTT GAA GTC CTT TTT
gly phe arg cys pro met asp cys leu tyr	gly leu gly lys thr leu glu val leu phe
62/21	92/31
ATT GGC AAT GCT GGA AAT GGA CAT TCC AAT	
ile gly asn ala gly asn gly his ser asn	ile ala arg ile asn arg thr arg OPA gly
120, 11	152/51
GGG GGC AAG CGT TTG TAC CGG GGC CAG CAA	
gly gly lys arg leu tyr arg gly gln gln	
	212/71
TGT TGT TGT GTC AGC GCG GGC TTG GTC TCG	
cys cys cys val ser ala gly leu val ser	
242/81 CAA AAC AGG TTG AAC TTA ACG ACT CAA GAA	272/91
gln asn arg leu asn leu thr thr gln glu	
-	332/111
302/101 GAC ACC AAT TTG ACT CGG CTC TTT GGC AAT	
asp thr asn leu thr arg leu phe gly asn	
362/121	392/131
GCA TCG TTG GCC TTG CCA TCA ATC GCC GGC	
ala ser leu ala leu pro ser ile ala gly	
422/141	452/151
GGA CCG ACC TCG ACC AGG GGT CCT TTG TGA	CTG CCG GGC TTG ACG CGG ACG ACC ACA GAG
gly pro thr ser thr arg gly pro leu OPA	leu pro gly leu thr arg thr thr thr glu
482/161	512/171
TCG GTC ATC GCC TAA GGC TAC CGT TCT GAC	
ser val ile ala OCH gly tyr arg ser asp	
542/181	572/191
GCA CGT CAT GTC TCA GCG GCC CAC CGC CAC	
ala arg his val ser ala ala his arg his	
602/201	632/211
AGA TGA CTC CAC GCA GCC TTG TTC GCA TCG	TTG GTG TCG TGG TTG CGA CGA CCT TGG CGC
arg OPA leu his ala ala leu phe ala ser	1eu vai ser trp leu arg arg pro trp arg 692/231
662/221	
TGG TGA GCG CAC CCG CCG GCG GTC GTG CCG trp OPA ala his pro pro ala val val pro	
trp OPA ata his pro pro ata vai vai pro	ary mee ary tre

SEQ ID No.13B

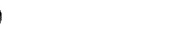
FIGURE 13B



									33/1									
GAT TTC																		
asp phe	val	ala	arg	trp	ile	val	cys	thr			glu	lys	his	leu	lys	ser	phe	leu
63/21									93/3									
TTG GCA																		
leu ala	met	leu	glu	met	asp	ile	pro	ile			glu	Leu	thr	giu	his	дтĀ	glu	дтА
123/41									153/		~~~	~~~		a	~~~	7.00	~~~	~~ T
GGG GCA																		
gly ala	ser	val	cys	thr	дтА	a⊥a	ser	lys			arg	pro	val	asp	arg	ser	gın	nıs
183/61 GTT GTT	am a	m ~ 3	666	acc	CCIII	mcc.	mcm	CCA	213/		ccc	CIDIT	ccc	TICC	N.C.C	ccc	mmC	mm.c
val val	vaı	ser	ата	arg	ala	crp	ser	ary	273/		gry	ieu	gry	crp	CIII	ary	Pire	pne
243/81 AAA ACA	CCT	ም <i>ር</i> አ	λ C·Π	ጥ አ አ	CCA	כיייכ	NAC	A A C	,		CCT	тсδ	מככ	GCG	ACG	ፐርር	כייר	cee
lys thr																		
303/101	gry	OLA	CIII	OC11	arg	ıcu	- y 5	4511	333/		414	0111	0112	~_~	· · · · ·	201		429
ACA CCA	יייד∆	TGA	СТС	GGC	тст	ттG	GCA	ATT			GAG	CTG	CGA	GCA	GCC	GGG	TGA	CCG
thr pro																		
363/121		0211		5-1					393/		J		,			, 1		-
CAT CGT	TGG	CCT	TGC	CAT	CAA	TCG	CCG	GCT	CGC	GGA	CGT	AGA	TAA	TCA	GCT	CAC	CGT	TGG
his arg																		
423/141	-	-	_		_				453/									
GAC CGA																		
asp arg	pro	arg	pro	gly	val	leu	cys	asp	cys	arg	ala	OPA	arg	gly	arg	pro	gln	ser
483/161									-	171								
CGG TCA																		
arg ser	ser	pro	lys	ala	thr	val	leu	thr			cys	val	gly	ala	asp	asp	val	arg
543/181									,	/191								~~-
CAC GTC																		
his val	met	ser	gln	arg	pro	thr	ala	thr			ala	дтЛ	ser	met	ser	ala	cys	ата
603/201										/211	~~	0.00	maa	C3 C	a. a	amm.	666	a am
GAT GAC																		
asp asp	ser	thr	gin	pro	cys	ser	nıs	arg			arg	дту	cys	asp	asp	Leu	gry	ala
663/221 GGT GAG	ccc	700	ccc	ccc	CCC	ሞሮር	TCC	ccc		/231	GGA	ጥር						
gly glu												10						
gry gra	arq	LHL	arg	ату	ary	2CT	~y.s	$a_{\perp}y$	$a \perp a$	Cys	$\rightarrow \rightarrow Y$							

SEQ ID No.13C

FIGURE 13C





part of the nucleotide sequence of seq13A 1/1 GGG TCC TTT GTG ACT GCC GGG CTT GAC GCG GAC GAC CAC AGA GTC GGT CAT CGC CTA AGG gly ser phe val thr ala gly leu asp ala asp asp his arg val gly his arg leu arg 91/31 61/21 CTA CCG TTC TGA CCT GGG GCT GCG TGG GCG CCG ACG ACG TGA GGC ACG TCA TGT CTC AGC leu pro phe OPA pro gly ala ala trp ala pro thr thr OPA gly thr ser cys leu ser 151/51 GGC CCA CCG CCA CCT CGG TCG CCG GCA GTA TGT CAG CAT GTG CAG ATG ACT CCA CGC AGC gly pro pro pro pro arg ser pro ala val cys gln his val gln met thr pro arg ser 211/71 CTT GTT CGC ATC GTT GGT GTC GTG GTT GCG ACC TTG GCG CTG GTG AGC GCA CCC GCC leu val arg ile val gly val val val ala thr thr leu ala leu val ser ala pro ala 241/81 GGC GGT CGT GCC GCG CAT GCG GAT C gly gly arg ala ala his ala asp

SEQ ID No.13A'

FIGURE 13A'

31/11 GGT CCT TTG TGA CTG CCG GGC TTG ACG CGG ACG ACC ACA GAG TCG GTC ATC GCC TAA GGC gly pro leu OPA leu pro gly leu thr arg thr thr thr glu ser val ile ala OCH gly 61/21 91/31 TAC CGT TCT GAC CTG GGG CTG CGT GGG CGC CGA CGT GAG GCA CGT CAT GTC TCA GCG tyr arg ser asp leu gly leu arg gly arg arg arg glu ala arg his val ser ala 151/51 121/41 GCC CAC CGC CAC CTC GGT CGC CGG CAG TAT GTC AGC ATG TGC AGA TGA CTC CAC GCA GCC ala his arg his leu gly arg arg gln tyr val ser met cys arg OPA leu his ala ala 211/71 TTG TTC GCA TCG TTG GTG TCG TGG TTG CGA CGA CCT TGG CGC TGG TGA GCG CAC CCG CCG leu phe ala ser leu val ser trp leu arg arg pro trp arg trp OPA ala his pro pro GCG GTC GTG CCG CGC ATG CGG ATC ala val val pro arg met arg ile

SEQ ID No.13B'

FIGURE 13B'

31/11 GTC CTT TGT GAC TGC CGG GCT TGA CGC GGA CGA CCA CAG AGT CGG TCA TCG CCT AAG GCT val leu cys asp cys arg ala OPA arg gly arg pro gln ser arg ser ser pro lys ala 91/31 61/21ACC GTT CTG ACC TGG GGC TGC GTG GGC GCC GAC GTG AGG CAC GTC ATG TCT CAG CGG thr val leu thr trp gly cys val gly ala asp asp val arg his val met ser gln arg 151/51 121/41 CCC ACC GCC ACC TCG GTC GCC GGC AGT ATG TCA GCA TGT GCA GAT GAC TCC ACG CAG CCT pro thr ala thr ser val ala gly ser met ser ala cys ala asp asp ser thr gln pro 211/71 181/61 TGT TCG CAT CGT TGG TGT CGT GGT TGC GAC GAC CTT GGC GCT GGT GAG CGC ACC CGC CGG cys ser his arg trp cys arg gly cys asp asp leu gly ala gly glu arg thr arg arg 241/81 CGG TCG TGC CGC GCA TGC GGA TC arg ser cys arg ala cys gly

> SEQ ID No.13C' FIGURE 13C'



sequence Rv1984c predicted by Cole et al. (Nature 393:537-544) and containing seq13A' 1/1 31/11atq act cca cgc agc ctt gtt cgc atc gtt ggt gtc gtg gtt gcg acg acc ttg gcg ctg Met thr pro arg ser leu val arg ile val gly val val ala thr thr leu ala leu 91/31 61/21 gtg agc gca ccc gcc ggc ggt cgt gcc gcg cat gcg gat ccg tgt tcg gac atc gcg gtc val ser ala pro ala gly gly arg ala ala his ala asp pro cys ser asp ile ala val 151/51 gtt tte get ege gge aeg eat eag get tet ggt ett gge gae gte ggt gag geg tte gte val phe ala arg gly thr his gln ala ser gly leu gly asp val gly glu ala phe val 211/71 181/61 gac tog ott acc tog caa gtt ggc ggg cgg tog att ggg gtc tac gcg gtg aac tac coa asp ser leu thr ser gln val gly gly arg ser ile gly val tyr ala val asn tyr pro 271/91 241/81 gca age gae gae tae ege geg age geg tea aae ggt tee gat gat geg age gee eae ate ala ser asp asp tyr arg ala ser ala ser asn gly ser asp asp ala ser ala his ile 331/111 cag ege ace gte gee age tge eeg aac ace agg att gtg ett ggt gge tat teg eag ggt gln arg thr val ala ser cys pro asn thr arg ile val leu gly gly tyr ser gln gly 391/131 qcq acq qtc atc gat ttg tcc acc tcg gcg atg ccg ccc gcg gtg gca gat cat gtc gcc ala thr val ile asp leu ser thr ser ala met pro pro ala val ala asp his val ala 451/151 421/141 get gte gee ett tte gge gag eea tee agt ggt tte tee age atg ttg tgg gge gge ggg ala val ala leu phe gly glu pro ser ser gly phe ser ser met leu trp gly gly gly 511/171 481/161 tog ttg cog aca ato ggt cog ctg tat ago tot aag aco ata aac ttg tgt got coo gac ser leu pro thr ile gly pro leu tyr ser ser lys thr ile asn leu cys ala pro asp 571/191 541/181 gat cca ata tgc acc gga ggc ggc aat att atg gcg cat gtt tcg tat gtt cag tcg ggg asp pro ile cys thr gly gly gly asn ile met ala his val ser tyr val gln ser gly 601/201 631/211 atg aca age cag geg geg aca tte geg geg aac agg ete gat cae gee gga tga met thr ser gln ala ala thr phe ala ala asn arg leu asp his ala gly OPA

SEQ ID No.13D

FIGURE 13D



Seq13F: ORF predicted by Cole et al. (Nature 393:537-544) and containing Rv1984c 31/11 1/1 tga ggc acg tca tgt ctc agc ggc cca ccg cca cct cgg tcg ccg gca gta tgt cag cat OPA gly thr ser cys leu ser gly pro pro pro pro arg ser pro ala val cys gln his 91/31 gtg cag atg act cca cgc agc ctt gtt cgc atc gtt ggt gtc gtg gtt gcg acg acc ttg val gln met thr pro arg ser leu val arg ile val gly val val val ala thr thr leu 151/51 121/41 gcg ctg gtg agc gca ccc gcc ggc ggt cgt gcc gcg cat gcg gat ccg tgt tcg gac atc ala leu val ser ala pro ala gly gly arg ala ala his ala asp pro cys ser asp ile 211/71 gcg gtc gtt ttc gct cgc ggc acg cat cag gct tct ggt ctt ggc gac gtc ggt gag gcg ala val val phe ala arg gly thr his gln ala ser gly leu gly asp val gly glu ala 271/91 241/81 ttc gtc gac tcg ctt acc tcg caa gtt ggc ggg cgg tcg att ggg gtc tac gcg gtg aac phe val asp ser leu thr ser gln val gly gly arg ser ile gly val tyr ala val asn 331/111 301/101 tac cca gca agc gac gac tac cgc gcg agc gcg tca aac ggt tcc gat gat gcg agc gcc tyr pro ala ser asp asp tyr arg ala ser ala ser asn gly ser asp asp ala ser ala 391/131 361/121 cac atc cag cgc acc gtc gcc agc tgc ccg aac acc agg att gtg ctt ggt ggc tat tcg his ile gln arg thr val ala ser cys pro asn thr arg ile val leu gly gly tyr ser 451/151 cag ggt gcg acg gtc atc gat ttg tcc acc tcg gcg atg ccg ccc gcg gtg gca gat cat gln gly ala thr val ile asp leu ser thr ser ala met pro pro ala val ala asp his 511/171 481/161 gtc gcc gct gtc gcc ctt ttc ggc gag cca tcc agt ggt ttc tcc agc atg ttg tgg ggc val ala ala val ala leu phe gly glu pro ser ser gly phe ser ser met leu trp gly 571/191 541/181 ggc ggg tcg ttg ccg aca atc ggt ccg ctg tat agc tct aag acc ata aac ttg tgt gct gly gly ser leu pro thr ile gly pro leu tyr ser ser lys thr ile asn leu cys ala 631/211 601/201 ccc gac gat cca ata tgc acc gga ggc ggc aat att atg gcg cat gtt tcg tat gtt cag pro asp asp pro ile cys thr gly gly gly asn ile met ala his val ser tyr val gln 691/231 661/221 tcg ggg atg aca agc cag gcg gcg aca ttc gcg gcg aac agg ctc gat cac gcc gga tga ser gly met thr ser gln ala ala thr phe ala ala asn arg leu asp his ala gly OPA

SEQ ID No.13F

FIGURE 13F



31/11 CCA CCG GGG CTG GAG GGG CGA ATG TGC GCC GAA CGC CGT CGG CCA ACT TGG CCG CTG AGG pro pro gly leu glu gly arg met cys ala glu arg arg pro thr trp pro leu arg 91/31 61/21GCG GCT GAT CCC CTG GCC CGA GAC GGG GCA AGC CAA TAG CGG CTC CAT CGG GCT TTG CTG ala ala asp pro leu ala arg asp gly ala ser gln AMB arg leu his arg ala leu leu 151/51 121/41 GTA GCG GTT CGG CGG GAA CCG AGC GCC GAC GTT GTC GGT GCC CGG TGA TAT ATT GGG TCA val ala val arg arg glu pro ser ala asp val val gly ala arg OPA tyr ile gly ser 211/71 181/61 GAC GGG TAT GGC GGC GAC TGA GGT GAT CTG CGA CAC GCC GCC GCG GTG CTC GAG CCA GGC asp gly tyr gly gly asp OPA gly asp leu arg his ala ala ala val leu glu pro gly 271/91 TTA CGA CCA GGG AAT TTC GAA AAT GTT ATT CAG AAC ATC TTG TAT CTC TTC CTC CGT GCC leu arg pro gly asn phe glu asn val ile gln asn ile leu tyr leu phe leu arg ala 331/111 ACC CCC TAG GTG TAG TGT TTT CGA GTA CCG GCA GAT CCC AGT TCA CCA GTC TCA CCA GAT thr pro AMB val AMB cys phe arg val pro ala asp pro ser ser pro val ser pro asp

SEQ ID No.14A

FIGURE 14A

32/11 CAC CGG GGC TGG AGG GGC GAA TGT GCG CCG AAC GCC GTC GGC CAA CTT GGC CGC TGA GGG his arg gly trp arg gly glu cys ala pro asn ala val gly gln leu gly arg OPA gly 92/31 CGG CTG ATC CCC TGG CCC GAG ACG GGG CAA GCC AAT AGC GGC TCC ATC GGG CTT TGC TGG arg leu ile pro trp pro glu thr gly gln ala asn ser gly ser ile gly leu cys trp 152/51 TAG CGG TTC GGC GGG AAC CGA GCG CCG ACG TTG TCG GTG CCC GGT GAT ATA TTG GGT CAG AMB arg phe gly gly asn arg ala pro thr leu ser val pro gly asp ile leu gly gln 212/71 ACG GGT ATG GCG GCG ACT GAG GTG ATC TGC GAC ACG CCG CCG CGG TGC TCG AGC CAG GCT thr gly met ala ala thr glu val ile cys asp thr pro pro arg cys ser ser gln ala 272/91 242/81 TAC GAC CAG GGA ATT TCG AAA ATG TTA TTC AGA ACA TCT TGT ATC TCT TCC TCC GTG CCA tyr asp gln gly ile ser lys met leu phe arg thr ser cys ile ser ser ser val pro 332/111 302/101 CCC CCT AGG TGT AGT GTT TTC GAG TAC CGG CAG ATC CCA GTT CAC CAG TCT CAC CAG ATC pro pro arg cys ser val phe glu tyr arg gln ile pro val his gln ser his gln ile

SEQ ID No.14B

FIGURE 14B

33/11

ACC GGG GCT GGA GGG GCG AAT GTG CGC CGA ACG CCG TCG GCC AAC TTG GCC GCT GAG GGC thr gly ala gly gly ala asn val arg arg thr pro ser ala asn leu ala ala glu gly 93/31 GGC TGA TCC CCT GGC CCG AGA CGG GGC AAG CCA ATA GCG GCT CCA TCG GGC TTT GCT GGT gly OPA ser pro gly pro arg arg gly lys pro ile ala ala pro ser gly phe ala gly 153/51 AGC GGT TCG GCG GGA ACC GAG CGC CGA CGT TGT CGG TGC CCG GTG ATA TAT TGG GTC AGA ser gly ser ala gly thr glu arg arg cys arg cys pro val ile tyr trp val arg 213/71 CGG GTA TGG CGG CGA CTG AGG TGA TCT GCG ACA CGC CGC CGC GGT GCT CGA GCC AGG CTT arg val trp arg arg leu arg OPA ser ala thr arg arg gly ala arg ala arg leu 273/91 243/81 ACG ACC AGG GAA TTT CGA AAA TGT TAT TCA GAA CAT CTT GTA TCT CTT CCT CCG TGC CAC thr thr arg glu phe arg lys cys tyr ser glu his leu val ser leu pro pro cys his 333/111 303/101 CCC CTA GGT GTA GTG TTT TCG AGT ACC GGC AGA TCC CAG TTC ACC AGT CTC ACC AGA TC pro leu gly val val phe ser ser thr gly arg ser gln phe thr ser leu thr arg

SEQ ID No.14C

FIGURE 14C

part of the nucleotide sequence of seq14A

1/1 \$31/11 TTT TCG AGT ACC GGC AGA TCC CAG GTT CAC CAG GTC TCA CCA GAT C phe ser ser thr gly arg ser gln val his gln val ser pro asp

SEQ ID No.14A'

FIGURE 14A'

1/1 31/11 TGT TTT CGA GTA CCG GCA GAT CCC AGG TTC ACC AGG TCT CAC CAG ATC cys phe arg val pro ala asp pro arg phe thr arg ser his gln ile

SEQ ID No.14C

FIGURE 14C

1/1 \$31/11\$ GTT TTC GAG TAC CGG CAG ATC CCA GGT TCA CCA GGT CTC ACC AGA TC val phe glu tyr arg gln ile pro gly ser pro gly leu thr arg

SEQ ID No.14C'

FIGURE 14C'

REPLACEMENT SHEET (RULE 26)



ORF predicted based on the sequence published by Cole et al. (Nature 393:537-544) and containing seq14A'

```
31/11
1/1
TAG CGG TTC GGC GGG AAG CTA GCG GCG ACG TTG TCG GTG GCC GGT GAT ATA TTG GGT CAG
AMB arg phe gly gly lys leu ala ala thr leu ser val ala gly asp ile leu gly gln
                                        91/31
61/21
ACG GGT ATG GCG GCG GCT GAG GTG ATC TGC GAC ACG CCG CCG CGG TGC TCG AGC CAG GCT
thr gly met ala ala ala glu val ile cys asp thr pro pro arg cys ser ser gln ala
                                        151/51
121/41
TAC GAC CAG GGA ATT TCG AAA ATG TTA TTC AGA ACA TCT TGT ATC TCT TCT CCG TGC CAC
tyr asp gln gly ile ser lys met leu phe arg thr ser cys ile ser ser pro cys his
                                        211/71
181/61
CCC CTA GGT GTA GTG TTT TCG AGT ACC GGC AGA TCC CAG GTT CAC CAG GTC TCA CCA gat
pro leu gly val val phe ser ser thr gly arg ser gln val his gln val ser pro asp
                                        271/91
241/81
cca cgg ggc gcg atg aac ttc ccg gca tcg gca tcg cca ggt cga cgg acg tgg tcg cgc
pro arg gly ala met asn phe pro ala ser ala ser pro gly arg arg thr trp ser arg
                                        331/111
301/101
tat gac ggg aat ctg gag cct tgt cgg gcc gct caa cat atc gaa gat gca cta ctt gag
tyr asp gly asn leu glu pro cys arg ala ala gln his ile glu asp ala leu leu glu
                                         391/131
tcg ttg cca gat cct gtc aga ttc ccg att tcc gca aag gag cgg tac gcc cat gac cgt
ser leu pro asp pro val arg phe pro ile ser ala lys glu arg tyr ala his asp arg
421/141
gac cgt tta cac taa
asp arg leu his OCH
```

SEQ ID No.14F

FIGURE 14F



Sequence Rv3054c predicted by Cole et al. (Nature 393:537-544) which may be in phase with Seq14A'

1/1	31/11
ofg toa gat acc aag too gac atc aaa atc	ttg gcc tta gtg gga agc ctg cgc gcg gcg
val ser asp thr lvs ser asp ile lys ile	e leu ala leu val gly ser leu arg ala ala
61/21	91/31
tog the aac ege cag ate gee gag etg get	ged aag gte get eeg gad gge gte ace gte
ser phe ash ard gln ile ala glu leu ala	ala lys val ala pro asp gly val thr val
121/41	151/51
acc ato tto gag ggg ctg ggg gac ctg ccg	tto tac aac gaa gac atc gac aca gcg acg
thr met phe glu glv leu glv asp leu pro	phe tyr asn glu asp ile asp thr ala thr
181/61	211/71
gan ata con aca con ata ago ded ttd ego	g gag gee geg tet gae geg cae get gee ttg
gly val pro ala pro val ser ala leu arg	g glu ala ala ser asp ala his ala ala leu
241/81	271/91
ata ata aca caa gaa tac aac agc agc att	ccg gcc gtg atc aag aac gcg atc gac tgg
val val thr pro glu tyr asn gly ser ile	e pro ala val ile lys asn ala ile asp trp
301/101	331/111
ctg too agg coa tto ggc gat ggc gcg tto	g aag gac aag ceg ttg gee gtg ate gge gge
leu ser arg pro phe gly asp gly ala len	lys asp lys pro leu ala val ile gly gly
361/121	391/131
tcc atg ggc cgc tac ggc ggg gta tgg gcc	g cac gac gag act cgc aag tcg ttc agc atc
ser met gly arg tyr gly gly val trp ala	a his asp glu thr arg lys ser phe ser ile
421/141	451/151
act and acq can atq atc gat gcg atc aa	a ctg tcg gtg ccg ttc caa act ctg ggc aag
ala gly thr arg val val asp ala ile ly	s leu ser val pro phe gln thr leu gly lys
481/161	511/171
ten ate aca dae dee dee dad eta ded de	g aat gtg cgc gac gcc gtc ggc aac ttg gcc
ser val ala asp asp ala gly leu ala al	a asn val arg asp ala val gly asn leu ala
541/181	
gct gag gtc ggc tga	
ala glu val gly OPA	

SEQ ID No.14R

FIGURE 14R



ORF predicted by Cole et al. (Nature 393:537-544) and containing Rv3054c

```
31/11
1/1
taa cgc gat cgg aat aaa tcg gac cat ggt ccg gtt ggc tcg tgc aag gac gtg gac caa
OCH arg asp arg asn lys ser asp his gly pro val gly ser cys lys asp val asp gln
                                        91/31
caa gcg gaa agg aac gta gca gtg tca gat acc aag tcc gac atc aaa atc ttg gcc tta
gln ala glu arg asn val ala val ser asp thr lys ser asp ile lys ile leu ala leu
                                        151/51
121/41
gtg gga age etg ege geg teg tte aae ege eag ate gee gag etg get gee aag gte
val gly ser leu arg ala ala ser phe asn arg gln ile ala glu leu ala ala lys val
                                        211/71
181/61
gct ccg gac ggc gtc acc gtc acc atg ttc gag ggg ctg ggg gac ctg ccg ttc tac aac
ala pro asp gly val thr val thr met phe glu gly leu gly asp leu pro phe tyr asn
                                        271/91
241/81
gaa gac atc gac aca gcg acg gag gtg ccg gcg ccg gtg agc gcg ttg cgg gag gcc gcg
glu asp ile asp thr ala thr glu val pro ala pro val ser ala leu arg glu ala ala
                                        331/111
tet gae geg cae get gee ttg gtg gte aeg eeg gaa tae aae gge age att eeg gee gtg
ser asp ala his ala ala leu val val thr pro glu tyr asn gly ser ile pro ala val
                                        391/131
361/121
atc aag aac gcg atc gac tgg ctg tcc agg cca ttc ggc gat ggc gcg ttg aag gac aag
ile lys asn ala ile asp trp leu ser arg pro phe gly asp gly ala leu lys asp lys
                                         451/151
421/141
ccg ttg gcc gtg atc ggc ggc tcc atg ggc cgc tac ggc ggg gta tgg gcg cac gac gag
pro leu ala val ile gly gly ser met gly arg tyr gly gly val trp ala his asp glu
                                         511/171
481/161
act cgc aag tcg ttc agc atc gct ggc acg cgg gtg gtc gat gcg atc aaa ctg tcg gtg
thr arg lys ser phe ser ile ala gly thr arg val val asp ala ile lys leu ser val
                                         571/191
541/181
ceg ttc caa act ctg ggc aag tcg gtc gcg gac gcc ggg ctg gcg gcg aat gtg cgc
pro phe gln thr leu gly lys ser val ala asp asp ala gly leu ala ala asn val arg
                                         631/211
601/201
gac gcc gtc ggc aac ttg gcc gct gag gtc ggc tga
asp ala val gly asn leu ala ala glu val gly OPA
```

SEQ ID No.14P

FIGURE 14P



fragment based on the sequence published by Cole et al. (Nature 393:537-544) and containing seq 14F' and seq 14P'

31/11 1/1 taa cgc gat cgg aat aaa tcg gac cat ggt ccg gtt ggc tcg tgc aag gac gtg gac caa OCH arg asp arg asn lys ser asp his gly pro val gly ser cys lys asp val asp gln asn ala ile gly ile asn arg thr met val arg leu ala arg ala arg thr trp thr asn thr arg ser glu OCH ile gly pro trp ser gly trp leu val gln gly arg gly pro thr 61/21 91/31 caa gcg gaa agg aac gta gca gtg tca gat acc aag tcc gac atc aaa atc ttg gcc tta gln ala glu arg asn val ala val ser asp thr lys ser asp ile lys ile leu ala leu lys arg lys gly thr AMB gln cys gln ile pro ser pro thr ser lys ser trp pro AMB ser gly lys glu arg ser ser val arg tyr gln val arg his gln asn leu gly leu ser 151/51 121/41 gtg gga agc ctg cgc gcg gcg tcg ttc aac cgc cag atc gcc gag ctg gct gcc aag gtc val gly ser leu arg ala ala ser phe asn arg gln ile ala glu leu ala ala lys val trp glu ala cys ala arg arg ser thr ala arg ser pro ser trp leu pro arg ser gly lys pro ala arg gly val val gln pro pro asp arg arg ala gly cys gln gly arg 211/71 181/61 gct ccg gac ggc gtc acc gtc acc atg ttc gag ggg ctg ggg gac ctg ccg ttc tac aac ala pro asp gly val thr val thr met phe glu gly leu gly asp leu pro phe tyr asn leu arg thr ala ser pro ser pro cys ser arg gly trp gly thr cys arg ser thr thr ser gly arg arg his arg his his val arg gly ala gly gly pro ala val leu gln arg 241/81 271/91 gaa gac atc gac aca gcg acg gag gtg ccg gcg ccg gtg agc gcg ttg cgg gag gcc gcg glu asp ile asp thr ala thr glu val pro ala pro val ser ala leu arg glu ala ala lys thr ser thr gln arg arg arg cys arg arg oPA ala arg cys gly arg pro arg arg his arg his ser asp gly gly ala gly ala gly glu arg val ala gly gly arg val 301/101 331/111 tet gac geg cac get gee ttg gtg gtc acg ccg gaa tac aac ggc agc att ccg gcc gtg ser asp ala his ala ala leu val val thr pro glu tyr asn gly ser ile pro ala val leu thr arg thr leu pro trp trp ser arg arg asn thr thr ala ala phe arg pro OPA OPA arg ala arg cys leu gly gly his ala gly ile gln arg gln his ser gly arg asp 361/121 391/131 atc aag aac gcg atc gac tgg ctg tcc agg cca ttc ggc gat ggc gcg ttg aag gac aag ile lys asn ala ile asp trp leu ser arg pro phe gly asp gly ala leu lys asp lys ser arg thr arg ser thr gly cys pro gly his ser ala met ala arg OPA arg thr ser gln glu arg asp arg leu ala val gln ala ile arg arg trp arg val glu gly gln ala 421/141 451/151 ccg ttg gcc gtg atc ggc ggc tcc atg ggc cgc tac ggc ggg gta tgg gcg cac gac gag pro leu ala val ile gly gly ser met gly arg tyr gly gly val trp ala his asp glu arg trp pro OPA ser ala ala pro trp ala ala thr ala gly tyr gly arg thr thr arg val gly arg asp arg leu his gly pro leu arg arg gly met gly ala arg arg asp 481/161 511/171 act cgc aag tcg ttc agc atc gct ggc acg cgg gtg gtc gat gcg atc aaa ctg tcg gtg thr arg lys ser phe ser ile ala gly thr arg val val asp ala ile lys leu ser val leu ala ser arg ser ala ser leu ala arg gly trp ser met arg ser asn cys arg cys ser gln val val gln his arg trp his ala gly gly arg cys asp gln thr val gly ala 541/181 571/191 ecg tte caa act etg gge aag teg gte geg gae gae gee ggg etg geg aat gtg ege pro phe gln thr leu gly lys ser val ala asp asp ala gly leu ala ala asn val arg arg ser lys leu trp ala ser arg ser arg thr thr pro gly trp arg arg met cys ala val pro asn ser gly gln val gly arg gly arg arg arg ala gly gly glu cys ala arg 631/211 601/201 gac gcc gtc ggc aac ttg gcc gct gag gtc ggc tga tcc ctg ggc cga ggc ggg tca gcc asp ala val gly asn leu ala ala glu val gly OPA ser leu gly arg gly gly ser ala thr pro ser ala thr trp pro leu arg ser ala asp pro trp ala glu ala gly gln pro arg arg gln leu gly arg OPA gly arg leu ile pro gly pro arg arg val ser gln

SEQ ID No.14Q

691/231

aat agc ggc tcc atc ggc ttt gct ggt agc ggt tcg gcg gga agc tag cgg cga cgt tgt asn ser gly ser ile gly phe ala gly ser gly ser ala gly ser AMB arg arg cys ile ala ala pro ser ala leu leu val ala val arg arg glu ala ser gly asp val val AMB arg leu his arg leu cys trp AMB arg phe gly gly lys leu ala ala thr leu ser

FIGURE 14Q



751/251 721/241 cgg tgg ccg gtg ata tat tgg gtc aga cgg gta tgg cgg cgg ctg agg tga tct gcg aca arg trp pro val ile tyr trp val arg arg val trp arg arg leu arg OPA ser ala thr gly gly arg OPA tyr ile gly ser asp gly tyr gly gly GPA gly asp leu arg his val ala gly asp ile leu gly gln thr gly met ala ala glu val ile cys asp thr 811/271 781/261 cgc cgc cgc ggt gct cga gcc agg ctt acg acc agg gaa ttt cga aaa tgt tat tca gaa arg arg gly ala arg ala arg leu thr thr arg glu phe arg lys cys tyr ser glu ala ala ala val leu glu pro gly leu arg pro gly asn phe glu asn val ile gln asn pro pro arg cys ser ser gln ala tyr asp gln gly ile ser lys met leu phe arg thr 871/291 841/281 cat ctt gta tct ctt ctc cgt gcc acc ccc tag gtg tag tgt ttt cga gta ccg gca gat his leu val ser leu leu arg ala thr pro AMB val AMB cys phe arg val pro ala asp ile leu tyr leu phe ser val pro pro pro arg cys ser val phe glu tyr arg gln ile ser cys ile ser ser pro cys his pro leu gly val val phe ser ser thr gly arg ser 931/311 901/301 ccc agg ttc acc agg tct cac cag atc cac ggg gcg cga tga act tcc cgg cat cgg cat pro arg phe thr arg ser his gln ile his gly ala arg OPA thr ser arg his arg his pro gly ser pro gly leu thr arg ser thr gly arg asp glu leu pro gly ile gly ile gln val his gln val ser pro asp pro arg gly ala met asn phe pro ala ser ala ser 991/331 961/321 cgc cag gtc gac gga cgt ggt cgc gct atg acg gga atc tgg agc ctt gtc ggg ccg ctc arg gln val asp gly arg gly arg ala met thr gly ile trp ser leu val gly pro leu ala arg ser thr asp val val ala leu OPA arg glu ser gly ala leu ser gly arg ser pro gly arg arg thr trp ser arg tyr asp gly asn leu glu pro cys arg ala ala gln 1051/351 1021/341 aac ata tog aag atg cao tao ttg agt ogt tgo cag atc otg toa gat too oga ttt oog asn ile ser lys met his tyr leu ser arg cys gln ile leu ser asp ser arg phe pro thr tyr arg arg cys thr thr OPA val val ala arg ser cys gln ile pro asp phe arg his ile glu asp ala leu leu glu ser leu pro asp pro val arg phe pro ile ser ala 1111/371 1081/361 caa agg agc ggt acg ccc atg acc gtg acc gtt tac act aa gln arg ser gly thr pro met thr val thr val tyr thr lys gly ala val arg pro OPA pro OPA pro phe thr leu lys glu arg tyr ala his asp arg asp arg leu his OCH

SEQ ID No.14Q(continued)

FIGURE 14Q (continued)

1 /1					31/11								
1/1 CAA GCC CGG CCG	CGA CTG	TTT GCC	GTT	TTG	GGG CTC	CTA	CCA	GAA	CAC	CAC	CTG	GCG	GCC
gln ala arg pro	ara lou	nhe ala	val	len	alv leu	leu	pro	alu	his	his	leu	ala	ala
-	arg reu	pire are		200	91/31		-	-					
61/21 GCG CAC CAT GGT	CILC CAC	ሮ እሮ ምጥረ	· cca	ጥሮር		CCC	GCG	CGC	GGG	CGG	CGA	CGA	CGT
GCG CAC CAT GGT	GTG CAC	CAG III	CGA	100	GII CCI	000	1.		-1	~~~	2 2 2	2 2 7	ara
ala his his gly	val his	gln le	ıarg	ser	val pro	pro	ата	arg	дтĀ	arg	ary	ary	arg
121/41		_			151/51								
CGA TGC CCG CGC	222 222	ccc cc	CCT	CCC	ጥልር ርጥር	GAC	CCG	GTC	GAC	GAC	GAC	GGG	GTC
CGA TGC CCG CGC	CCC GGC	GGC GCA	r GCI	900	ING CIC			*** 1	267	267	aen	alv	val
arg cys pro arg	pro gly	gly ala	a ala	ala	AMB Leu	asp	bro	vai	asp	asp	asp	9+1	
101/61					211//1								
GGC GGA CCA GTC	CCC GAT	GTC GAG	GCG	ATG	GCA ATA	CAG	CGC	CTT	GGT	GCG	CGG	ÇCA	CAC
GGC GGA CCA GIC	GGC GAI	010 021	1 -		-11-	aln	ara	1 611	alv	ala	ard	pro	his
gly gly pro val	gly asp	val gi	ı aıa	met	ara ire	gin	ary	rcu	9+1	ara	~-9	F	
241/81					271/91								
GTC TGA GGT GGC	GAA GAC	CAG TC	CGC	GCC	CAC CGG	CAG	CCG	GAT	CCG	GAT	ACG	CGG	TAC
val OPA gly gly	JAN GAO	-1		212	hie ara	aln	nro	asp	pro	asp	thr	arq	tyr
val OPA gly gly	gru asp	gin se	Larg	ата	mrs arg	9111	PLO	~DP	r = 0				-

SEQ ID No.15A

FIGURE 15A



32/11 AAG CCC GGC CGC GAC TGT TTG CCG TTT TGG GGC TCC TAC CAG AAC ACC ACC TGG CGG CCG lys pro gly arg asp cys leu pro phe trp gly ser tyr gln asn thr thr trp arg pro 92/31 62/21 CGC ACC ATG GTG TGC ACC AGT TGC GAT CGG TTC CTC CCG CGC GCG GGC GAC GAC GTC arg thr met val cys thr ser cys asp arg phe leu pro arg ala gly gly asp asp val 152/51 122/41 GAT GCC CGC GCC CCG GCG GCG CAG CTG CGT AGC TCG ACC CGG TCG ACG ACG ACG GGG TCG asp ala arg ala pro ala ala gln leu arg ser ser thr arg ser thr thr thr gly ser 212/71 182/61 GCG GAC CAG TCG GCG ATG TCG AGG CGA TGG CAA TAC AGC GCC TTG GTG CGC GGC CAC ACG ala asp gln ser ala met ser arg arg trp gln tyr ser ala leu val arg gly his thr 272/91 242/81 TCT GAG GTG GCG AAG ACC AGT CCC GCG CCC ACC GGC AGC CGG ATC CGG ATA CGC GGT AC ser glu val ala lys thr ser pro ala pro thr gly ser arg ile arg ile arg gly

SEQ ID No.15B

FIGURE 15B

33/11 AGC CCG GCC GCG ACT GTT TGC CGT TTT GGG GCT CCT ACC AGA ACA CCA CCT GGC GGC CGC ser pro ala ala thr val cys arg phe gly ala pro thr arg thr pro pro gly gly arg 93/31 GCA CCA TGG TGT GCA CCA GTT GCG ATC GGT TCC TCC CGC GCG CGG GCG GCG ACG ACG TCG ala pro trp cys ala pro val ala ile gly ser ser arg ala arg ala ala thr thr ser 153/51 123/41 ATG CCC GCG CCC CGG CGC CGC AGC TGC GTA GCT CGA CCC GGT CGA CGA CGA CGG GGT CGG met pro ala pro arg arg arg ser cys val ala arg pro gly arg arg arg arg gly arg 213/71 183/61 CGG ACC AGT CGG CGA TGT CGA GGC GAT GGC AAT ACA GCG CCT TGG TGC GCG GCC ACA CGT arg thr ser arg arg cys arg gly asp gly asn thr ala pro trp cys ala ala thr arg 273/91 243/81 CTG AGG TGG CGA AGA CCA GTC CCG CGC CCA CCG GCA GCC GGA TCC GGA TAC GCG GTA C leu arg trp arg arg pro val pro arg pro pro ala ala gly ser gly tyr ala val

SEQ ID No.15C

FIGURE 15C





part of the nucleotide sequence of seq15A

SEQ ID No.15A'

FIGURE 15A'

1/1			31/11	
GCG GCC GCG CGC				CCG CGC GCG GGC GAC
ala ala ala arg	his gly val	his gln leu	arg ser val leu p	oro arg ala gly gly asp
61/21			91/31	
				GAC CCG GTC GAC GAC GAC
asp val asp gly	arg ala pro	ala ala ala	ala ala AMB leu a	asp pro val asp asp asp
121/41			151/51	
				CAG CGC CTT GGT GCG CGG
gly val gly gly	pro val gly	asp val glu	ala met ala ile	gln arg leu gly ala arg
181/61			211/71	
			CGC GCC CAC CGG	
pro his val OPA	gly gly glu	asp gln ser	arg ala his arg	gln pro asp

SEQ ID No.15B'

FIGURE 15B'

1/1		31/11	
TGG CGG CCG CGC	GCC ATG GTG TGC ACC AGT	TGC GAT CGG TTC TCC CGC G	GCG CGG GCG GCG
trp arg pro arg	ala met val cys thr ser	cys asp arg phe ser arg a	ala arg ala ala
61/21		91/31	
		CAG CTG CGT AGC TCG ACC	
thr thr ser met	ala ala pro arg arg leu	gln leu arg ser ser thr a	arg ser thr thr
121/41		151/51	
ACG GGG TCG GCG	GGC CAG TCG GCG ATG TCG	AGG CGA TGG CAA TAC AGC	SCC TTG GTG CGC
thr gly ser ala	gly gln ser ala met ser	arg arg trp gln tyr ser a	ala leu val arg
181/61		211/71	
GGC CAC ACG TCT	GAG GTG GCG AAG ACC AGT	CCC GCG CCC ACC GGC AGC C	CGG ATC
gly his thr ser	glu val ala lys thr ser	pro ala pro thr gly ser a	arg ile

SEQ ID No.15C'

FIGURE 15C'

REPLACEMENT SHEET (RULE 26)



ORF containing Seq15A' according to Cole et al. (Nature 393:537-544)

1/1	31/11
taa ggt ccg cca acg ctt tac gct cga c	egg eeg eea ega gtt gge egg eea ett tea gge
OCH gly pro pro thr leu tyr ala arg a	arg pro pro arg val gly arg pro leu ser gly
61/21	91/31
cgt agt cgc cgc agg gca ggg ctt ccc g	gcg tcg tct tcg cgg gtt tgt cgg caa agg tgt
arg ser arg arg ala gly leu pro a	ala ser ser ser arg val cys arg gln arg cys
121/41	151/51
agg ggt agc gtt cgt ggg cgt cga cga c	ega tgt gca gct cgg gga tgc cgg cgc ggg
arg gly ser val arg gly arg arg arg	arg cys ala ala arg gly cys arg arg gly
181/61	211/71
cgg tgg ggg tgc gca cgc ccg gcc gcg a	act gtt tgc gcg ttt tgg ggc tct gcc aga aca
arg trp gly cys ala arg pro ala ala t	thr val cys ala phe trp gly ser ala arg thr
241/81	271/91
cca cct ggc ggc cgc gcg cca tgg tgt g	gca cca gtt gcg atc ggt tct ccc gcg cgc ggg
pro pro gly gly arg ala pro trp cys a	ala pro val ala ile gly ser pro ala arg gly
301/101	331/111
cgg cga cga cgt cga tgg ccg cgc ccc g	gge gge tge age tge gta get ega eee ggt ega
arg arg arg arg trp pro arg pro	gly gly cys ser cys val ala arg pro gly arg
361/121	391/131
cga cga cgg ggt cgg cgg gcc agt cgg c	cga tgt cga ggc gat ggc aat aca gcg cct tgg
arg arg arg gly arg arg ala ser arg a	arg cys arg gly asp gly asn thr ala pro trp
421/141	451/151
tgc gcg gcc aca cgt ctg agg tgg cga a	aga cca gtc ccg cgc cca ccg gca gcc gga tca
	arg pro val pro arg pro pro ala ala gly ser
481/161	511/171
ggt agg gca ggc gcg agt ctt cag cgg (ggt tgg cgg cga cga gca gct cca cag agt gtg
	gly trp arg arg ala ala pro gln ser val
541/181	571/191
agg gta cgg gcg gcg tac ggc aac ggt	gaa gca ggc act ccg acg aac cca tcg tca cgt
= -	glu ala gly thr pro thr asn pro ser ser arg
601/201	
cga agg ggc agg tga	
arg arg gly arg OPA	

SEQ ID No.15F

FIGURE 15F

R:Rv2530c predicted according to Cole et al. (Nature 393:537-544) and which may be in phase with SEQ15A 31/11 1/1 gtg acg gca ctg ctc gat gtc aat gtg ctg atc gcg ctg ggc tgg ccg aat cac gtt cac val thr ala leu leu asp val asn val leu ile ala leu gly trp pro asn his val his 91/31 cat gcg gcc gcg cag cga tgg ttc acg cag ttc tcc tcg aat ggg tgg gcc acc acg ccg his ala ala ala gln arg trp phe thr gln phe ser ser asn gly trp ala thr thr pro 151/51 121/41 atc acc qaq qca qqq tat qtc cga att tca agc aat cgc agt gtg atg cag gtg tcg acc ile thr glu ala gly tyr val arg ile ser ser asn arg ser val met gln val ser thr 211/71 acg ccg gct atc gcg atc gct cag ttg gcg gcg atg act tct ctt gcc ggg cac acg ttt thr pro ala ile ala ile ala gln leu ala ala met thr ser leu ala gly his thr phe 271/91 241/81 tqq cct qac qat gtg cca ctg atc gtt ggg agc gcc ggc gat cgc gat gcg gtg tcc aac trp pro asp asp val pro leu ile val gly ser ala gly asp arg asp ala val ser asn 331/111 cac cgt cgg gtc acc gac tgc cat ctc atc gcc ttg gcc gcg cgc tac ggg ggc cgg ttg his arg arg val thr asp cys his leu ile ala leu ala arg tyr gly gly arg leu 391/131 361/121 qtc aca ttc gat gcc gca ctg gcc gat tca gca tcc gca ggc ctc gtc gag gtg ttg tag val thr phe asp ala ala leu ala asp ser ala ser ala gly leu val glu val leu AMB

SEQ ID No.15R

FIGURE 15R

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Seq15P: ORF according to Cole et al. (Nature 393:537-544) containing Rv2530c
1/1
                                        31/11
tga tgt too goo gga tgo goo gao ggt gao tto oga gga tgt ogt oog ogo got oga gga
OPA cys ser ala gly cys ala asp gly asp phe arg gly cys arg pro arg ala arg gly
                                        91/31
61/21
cga cgt gtg acg gca ctg ctc gat gtc aat gtg ctg atc gcg ctg ggc tgg ccg aat cac
arg arg val thr ala leu leu asp val asn val leu ile ala leu gly trp pro asn his
                                        151/51
121/41
gtt cac cat gcg gcc gcg cag cga tgg ttc acg cag ttc tcc tcg aat ggg tgg gcc acc
val his his ala ala ala gln arg trp phe thr gln phe ser ser asn gly trp ala thr
                                        211/71
181/61
acg ccg atc acc gag gca ggg tat gtc cga att tca agc aat cgc agt gtg atg cag gtg
thr pro ile thr glu ala gly tyr val arg ile ser ser asn arg ser val met gln val
                                        271/91
241/81
tcg acc acg ccg gct atc gcg atc gct cag ttg gcg gcg atg act tct ctt gcc ggg cac
ser thr thr pro ala ile ala ile ala gln leu ala ala met thr ser leu ala gly his
                                        331/111
acq ttt tqq cct gac gat gtg cca ctg atc gtt ggg agc gcc ggc gat cgc gat gcg gtg
thr phe trp pro asp asp val pro leu ile val gly ser ala gly asp arg asp ala val
                                        391/131
tcc aac cat cgg gtc acc gac tgc cat ctc atc gcc ttg gcc gcg cgc tac ggg ggc
ser asn his arg arg val thr asp cys his leu ile ala leu ala ala arg tyr gly
                                        451/151
cgg ttg gtc aca ttc gat gcc gca ctg gcc gat tca gca tcc gca ggc ctc gtc gag gtg
arg leu val thr phe asp ala ala leu ala asp ser ala ser ala gly leu val glu val
481/161
ttg tag
leu AMB
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SEQ ID No.15P

FIGURE 15P REPLACEMENT SHEET (RULE 26)



Fragment containing Seq15P' and Seq 15F' tga tgt tcc gcc gga tgc gcc gac ggt gac ttc cga gga tgt cgt ccg cgc gct cga gga OPA cys ser ala gly cys ala asp gly asp phe arg gly cys arg pro arg ala arg gly asp val pro pro asp ala pro thr val thr ser glu asp val val arg ala leu glu asp Met phe arg arg met arg arg oPA leu pro arg met ser ser ala arg ser arg thr 91/31 61/21 cga cgt gtg acg gca ctg ctc gat gtc aat gtg ctg atc gcg ctg ggc tgg ccg aat cac arg arg val thr ala leu leu asp val asn val leu ile ala leu gly trp pro asn his asp val OPA arg his cys ser met ser met cys OPA ser arg trp ala gly arg ile thr thr cys asp gly thr ala arg cys gln cys ala asp arg ala gly leu ala glu ser arg 151/51 121/41 gtt cac cat gcg gcc gcg cag cga tgg ttc acg cag ttc tcc tcg aat ggg tgg gcc acc val his his ala ala ala gln arg trp phe thr gln phe ser ser asn gly trp ala thr phe thr met arg pro arg ser asp gly ser arg ser ser pro arg met gly gly pro pro ser pro cys gly arg ala ala met val his ala val leu leu glu trp val gly his his 211/71 181/61 acg ccg atc acc gag gca ggg tat gtc cga att tca agc aat cgc agt gtg atg cag gtg thr pro ile thr glu ala gly tyr val arg ile ser ser asn arg ser val met gln val arg arg ser pro arg gln gly met ser glu phe gln ala ile ala val OPA cys arg cys ala asp his arg gly arg val cys pro asn phe lys gln ser gln cys asp ala gly val 271/91 241/81 tog acc acg ccg gct atc gcg atc gct cag ttg gcg gcg atg act tot ott gcc ggg cac ser thr thr pro ala ile ala ile ala gln leu ala ala met thr ser leu ala gly his arg pro arg arg leu ser arg ser leu ser trp arg arg OPA leu leu pro gly thr asp his ala gly tyr arg asp arg ser val gly gly asp asp phe ser cys arg ala his 331/111 301/101 acg ttt tgg cct gac gat gtg cca ctg atc gtt ggg agc gcc ggc gat cgc gat gcg gtg thr phe trp pro asp asp val pro leu ile val gly ser ala gly asp arg asp ala val arg phe gly leu thr met cys his OPA ser leu gly ala pro ala ile ala met arg cys val leu ala OPA arg cys ala thr asp arg trp glu arg arg arg ser arg cys gly val 391/131 361/121 tcc aac cac cgt cgg gtc acc gac tgc cat ctc atc gcc ttg gcc gcg cgc tac ggg ggc ser asn his arg arg val thr asp cys his leu ile ala leu ala ala arg tyr gly gly pro thr thr val gly ser pro thr ala ile ser ser pro trp pro arg ala thr gly ala gln pro pro ser gly his arg leu pro ser his arg leu gly arg ala leu arg gly pro 451/151 421/141 cgg ttg gtc aca ttc gat gcc gca ctg gcc gat tca gca tcc gca ggc ctc gtc gag gtg arg leu val thr phe asp ala ala leu ala asp ser ala ser ala gly leu val glu val gly trp ser his ser met pro his trp pro ile gln his pro gln ala ser ser arg cys val gly his ile arg cys arg thr gly arg phe ser ile arg arg pro arg arg gly val 511/171 481/161 ttg tag tca ccg ggg atg ggc tcg cca ggc ctg cag gat ctg cgg gcg cag gcg ccc leu AMB ser pro gly met gly gly ser pro gly leu gln asp leu arg ala gln ala pro cys ser his arg gly trp ala ala arg gln ala cys arg ile cys gly arg arg arg pro val val thr gly asp gly arg leu ala arg pro ala gly ser ala gly ala gly ala pro 571/191 541/181 ccg gtc gga cac cgg cag gcc gac gct ttt ggc cca cgc gcg cag ctc ggc gct gct ggg pro val gly his arg gln ala asp ala phe gly pro arg ala gln leu gly ala ala gly arg ser asp thr gly arg pro thr leu leu ala his ala arg ser ser ala leu leu gly gly arg thr pro ala gly arg arg phe trp pro thr arg ala ala arg arg cys trp ala 631/211 601/201 ctc ggg ctc ggc ggc agc cgg ctc gaa aac cgt ggt ggc gtc ggc atc gtc gac gaa cca leu gly leu gly gly ser arg leu glu asn arg gly gly val gly ile val asp glu pro ser gly ser ala ala gly ser lys thr val val ala ser ala ser ser thr asn gln arg ala arg arg gln pro ala arg lys pro trp trp arg arg his arg arg thr arg

SEQ ID No.15Q

FIGURE 150



691/231 661/221 ggt gag ggc ggc ggc tag ata gcg gta ggt gta ttc ctg ggc gag ctt gcg ggt ttg gca gly glu gly gly gly AMB ile ala val gly val phe leu gly glu leu ala gly leu ala val arg ala ala arg AMB arg AMB val tyr ser trp ala ser leu arg val trp gln OPA gly arg arg leu asp ser gly arg cys ile pro gly arg ala cys gly phe gly arg 721/241 751/251 gaa cac gat cgg cac gtt ggg aaa gcc gat ctg caa ttc ggc cag ccc atc ggc gat cgc glu his asp arg his val gly lys ala asp leu gln phe gly gln pro ile gly asp arg asn thr ile gly thr leu gly lys pro ile cys asn ser ala ser pro ser ala ile ala thr arg ser ala arg trp glu ser arg ser ala ile arg pro ala his arg arg ser pro 811/271 781/261 cgt cgg gcg ggc gaa gga gtg cgc gaa gat ctc cga gta gcg gtc ctc gac cac cac ggc arg arg ala gly glu gly val arg glu asp leu arg val ala val leu asp his his gly val gly arg ala lys glu cys ala lys ile ser glu AMB arg ser ser thr thr thr ala ser gly gly arg arg ser ala arg arg ser pro ser ser gly pro arg pro pro arg arg 871/291 841/281 ggc ccg tgg cag cgc ggc cag ttc ggt cag ttg gta ttt cag gtt gcc gtt cag cac gcc gly pro trp gln arg gly gln phe gly gln leu val phe gln val ala val gln his ala ala arg gly ser ala ala ser ser val ser trp tyr phe arg leu pro phe ser thr pro pro val ala ala arg pro val arg ser val gly ile ser gly cys arg ser ala arg gln 901/301 931/311 aga agt aag gtc cgc caa cgc ttt acg ctc gac ggc cgc cac gag ttg gcc ggc cac ttt arg ser lys val arg gln arg phe thr leu asp gly arg his glu leu ala gly his phe glu val arg ser ala asn ala leu arg ser thr ala ala thr ser trp pro ala thr phe lys OCH gly pro pro thr leu tyr ala arg pro pro arg val gly arg pro leu ser 991/331 961/321 cag gcc gta gtc gcc gca ggg cag ggc ttc ccg cgt cgt ctt cgc ggg ttt gtc ggc aaa gin ala val val ala ala gly gln gly phe pro arg arg leu arg gly phe val gly lys arg pro AMB ser pro gln gly arg ala ser arg val val phe ala gly leu ser ala lys gly arg ser arg arg arg ala gly leu pro ala ser ser ser arg val cys arg gln arg 1051/351 1021/341 ggt gta ggg gta gcg ttc gtg ggc gtc gac gat gtg cag ctc ggg gat gcc ggc ggc gly val gly val ala phe val gly val asp asp asp val gln leu gly asp ala gly gly val AMB gly AMB arg ser trp ala ser thr thr met cys ser ser gly met pro ala ala cys arg gly ser val arg gly arg arg arg cys ala ala arg gly cys arg arg arg 1111/371 1081/361 gcg ggc ggt ggg ggt gcg cac gcc cgg ccg cga ctg ttt gcg cgt ttt ggg gct ctg cca ala gly gly gly gly ala his ala arg pro arg leu phe ala arg phe gly ala leu pro arg ala val gly val arg thr pro gly arg asp cys leu arg val leu gly leu cys gln gly arg trp gly cys ala arg pro ala ala thr val cys ala phe trp gly ser ala arg 1171/391 1141/381 gaa cac cac ctg gcg gcc gcg cgc cat ggt gtg cac cag ttg cga tcg gtt ctc ccg cgc glu his his leu ala ala ala arg his gly val his gln leu arg ser val leu pro arg asn thr thr trp arg pro arg ala met val cys thr ser cys asp arg phe ser arg ala thr pro pro gly gly arg ala pro trp cys ala pro val ala ile gly ser pro ala arg 1231/411 1201/401 geg gge gge gae gae gte gat gge ege gee eeg geg get gea get geg tag ete gae eeg ala gly gly asp asp val asp gly arg ala pro ala ala ala ala ala AMB leu asp pro arg ala ala thr thr ser met ala ala pro arg arg leu gln leu arg ser ser thr arg gly arg arg arg arg trp pro arg pro gly gly cys ser cys val ala arg pro gly 1291/431 1261/421 gtc gac gac gac ggg gtc ggg gca gtc ggc gat gtc gag gcg atg gca ata cag cgc val asp asp asp gly val gly gly pro val gly asp val glu ala met ala ile gln arg ser thr thr thr gly ser ala gly gln ser ala met ser arg arg trp gln tyr ser ala arg arg arg gly arg arg ala ser arg arg cys arg gly asp gly asn thr ala pro

SEQ ID No.15Q (continued 1)

FIGURE 15Q (continued 1)

1351/451 1321/441 ctt ggt gcg cgg cca cac gtc tga ggt ggc gaa gac cag tcc cgc gcc cac cgg cag ccg leu gly ala arg pro his val OPA gly gly glu asp gln ser arg ala his arg gln pro leu val arg gly his thr ser glu val ala lys thr ser pro ala pro thr gly ser arg trp cys ala ala thr arg leu arg trp arg pro val pro arg pro pro ala ala gly 1411/471 1381/461 gat cag gta ggg cag gcg cga gtc ttc agc ggg gtt ggc ggc gac gag cag ctc cac aga asp gln val gly gln ala arg val phe ser gly val gly gly asp glu gln leu his arg ile arg AMB gly arg arg glu ser ser ala gly leu ala ala thr ser ser ser thr glu ser gly arg ala gly ala ser leu gln arg gly trp arg arg arg ala ala pro gln ser 1471/491 1441/481 qtq tga ggg tac ggg cgg cgt acg gca acg gtg aag cag gca ctc cga cga acc cat cgt val OPA gly tyr gly arg arg thr ala thr val lys gln ala leu arg arg thr his arg cys glu gly thr gly gly val arg gln arg OPA ser arg his ser asp glu pro ile val val arg val arg ala ala tyr gly asn gly glu ala gly thr pro thr asn pro ser ser 1501/501 cac gtc gaa ggg gca ggt ga his val glu gly ala gly thr ser lys gly gln val arg arg gly arg OPA

SEQ ID No.15Q (continued 2)

FIGURE 15Q (continued (2)

31/11 TGC GCA TGC CGA CCA GTG TGG TTG GCC GGA GTT CGT TTG TTC GCG ATT GCC TCA ACG ATT cys ala cys arg pro val trp leu ala gly val arg leu phe ala ile ala ser thr ile 91/31 61/21 CGA TAT AAC CAC TCT AGT CAC ATC AAC CAC ACT CGT ACC ATC GAG CGT GTG GGT TCA TGC arg tyr asn his ser ser his ile asn his thr arg thr ile glu arg val gly ser cys 151/51 121/41 CAT GCA TTC GCG ACC GCG GGA GCC GGC GAA CCC GGC GCC ACA CAT AAT CCA GAT TGA GGA his ala phe ala thr ala gly ala gly glu pro gly ala thr his asn pro asp OPA gly 211/71 GAC TTC CGT GCC GAA CCG ACG CCG ACG CAA GCT TTC GAC AGC CAT GAG CGC GGT CGC CGC asp phe arg ala glu pro thr pro thr gln ala phe asp ser his glu arg gly arg arg 271/91 241/81 CCT GGC AGT TGC AAG TCC TTG TGC ATA TTT TCT TGT CTA CGA ATC AAC CGA AAC GAC CGA pro gly ser cys lys ser leu cys ile phe ser cys leu arg ile asn arg asn asp arg 331/111 GCG GCC CGA GCA CCA TGA ATT CAA GCA GGC GGC GGT GTT GAC CGA CCT GCC CGG CGA GCT ala ala arg ala pro OPA ile gln ala gly gly val asp arg pro ala arg arg ala 391/131 361/121 GAT GTC CGC GCT ATC GCA GGG GTT GTC CCA GTT CGG GAT C asp val arg ala ile ala gly val val pro val arg asp

SEQ ID No.16A

FIGURE 16A





57/185

32/11

GCG CAT GCC GAC CAG TGT GGT TGG CCG GAG TTC GTT TGT TCG CGA TTG CCT CAA CGA TTC ala his ala asp gln cys gly trp pro glu phe val cys ser arg leu pro gln arg phe 92/31 GAT ATA ACC ACT CTA GTC ACA TCA ACC ACA CTC GTA CCA TCG AGC GTG TGG GTT CAT GCC asp ile thr thr leu val thr ser thr thr leu val pro ser ser val trp val his ala 152/51 ATG CAT TCG CGA CCG CGG GAG CCG GCG AAC CCG GCG CCA CAC ATA ATC CAG ATT GAG GAG

met his ser arg pro arg glu pro ala asn pro ala pro his ile ile gln ile glu glu 212/71 ACT TCC GTG CCG AAC CGA CGC CGA CGC AAG CTT TCG ACA GCC ATG AGC GCG GTC GCC GCC

thr ser val pro asn arg arg arg lys leu ser thr ala met ser ala val ala ala 272/91 242/81 CTG GCA GTT GGA AGT CCT TGT GCA TAT TTT CTT GTC TAC GAA TCA ACC GAA ACG ACC GAG

leu ala val ala ser pro cys ala tyr phe leu val tyr glu ser thr glu thr thr glu 332/111 302/101

CGG CCC GAG CAC CAT GAA TTC AAG CAG GCG GCG GTG TTG ACC GAC CTG CCC GGC GAG CTG arg pro glu his his glu phe lys gln ala ala val leu thr asp leu pro gly glu leu 392/131

ATG TCC GCG CTA TCG CAG GGG TTG TCC CAG TCC GGG ATC met ser ala leu ser gln gly leu ser gln phe gly ile

cys pro arg tyr arg arg gly cys pro ser ser gly

SEQ ID No.16B

FIGURE 16B

33/11

CGC ATG CCG ACC AGT GTG GTT GGC CGG AGT TCG TTT GTT CGC GAT TGC CTC AAC GAT TCG arg met pro thr ser val val gly arg ser ser phe val arg asp cys leu asn asp ser 63/21 93/31 ATA TAA CCA CTC TAG TCA CAT CAA CCA CAC TCG TAC CAT CGA GCG TGT GGG TTC ATG CCA ile OCH pro leu AMB ser his gln pro his ser tyr his arg ala cys gly phe met pro 153/51 TGC ATT CGC GAC CGC GGG AGC CGG CGA ACC CGG CGC CAC ACA TAA TCC AGA TTG AGG AGA cys ile arg asp arg gly ser arg arg thr arg arg his thr OCH ser arg leu arg arg 213/71 183/61 CTT CCG TGC CGA ACC GAC GCC GAC GCA AGC TTT CGA CAG CCA TGA GCG CGG TCG CCC leu pro cys arg thr asp ala asp ala ser phe arg gln pro OPA ala arg ser pro pro 273/91 TGG CAG TTG CAA GTC CTT GTG CAT ATT TTC TTG TCT ACG AAT CAA CCG AAA CGA CCG AGC trp gln leu gln val leu val his ile phe leu ser thr asn gln pro lys arg pro ser 333/111 303/101 GGC CCG AGC ACC ATG AAT TCA AGC AGG CGG CGG TGT TGA CCG ACC TGC CCG GCG AGC TGA gly pro ser thr met asn ser ser arg arg cys OPA pro thr cys pro ala ser OPA 363/121 393/131 TGT CCG CGC TAT CGC AGG GGT TGT CCC AGT TCG GGA TC

SEQ ID No.16C

FIGURE 16C

31/11 GCG GGC CAC CGA TCA GTC GAT CGG GTG GTT TCC GCT CCA TCA GCC CGG AAT TGA GGT GCC ala gly his arg ser val asp arg val val ser ala pro ser ala arg asn OPA gly ala 91/31 61/21GCA GTG ACG ACA CCA GCG CAG GAC GCG CCG TTG GTG TTT CCC TCT GTT GCT TTC CCG TCC ala val thr thr pro ala gln asp ala pro leu val phe pro ser val ala phe pro ser 151/51 121/41 GGC TCG CCT TTT TTT CAT CAA CGT TGG ACT GCC GCA GTG GCG ATG TTG GTC GCC GGC GTG gly ser pro phe phe his gln arg trp thr ala ala val ala met leu val ala gly val 211/71 TTC GGT CAC CTG ACG GTC GGG ATG TTC CTT GGG TCT CGG GTT GCT GCT GGG TTT GCT CAA phe gly his leu thr val gly met phe leu gly ser arg val ala ala gly phe ala gln 271/91 TGC CCT GCT GGT GCG GCG TTC GGC CGA GTC GAT CAC CGC CAA AGA GCA CCC GTT AAA ACG cys pro ala gly ala ala phe gly arg val asp his arg gln arg ala pro val lys thr 331/111 301/101 GTC GAT GGC CCT CAA CTC GGC ATC GCG ACT GGC GAT TAT CAC CAT GCC TCG GGC TGA TC val asp gly pro gln leu gly ile ala thr gly asp tyr his his ala ser gly OPA

SEO ID No.17A

FIGURE 17A

32/11 CGG GCC ACC GAT CAG TCG ATC GGG TGG TTT CCG CTC CAT CAG CCC GGA ATT GAG GTG CCG arg ala thr asp gln ser ile gly trp phe pro leu his gln pro gly ile glu val pro 92/31 CAG TGA CGA CAC CAG CGC AGG ACG CGC CGT TGG TGT TTC CCT CTG TTG CTT TCC CGT CCG gln OPA arg his gln arg arg thr arg arg trp cys phe pro leu leu ser arg pro 152/51 122/41 GCT CGC CTT TTT TTC ATC AAC GTT GGA CTG CCG CAG TGG CGA TGT TGG TCG CCG GCG TGT ala arg leu phe phe ile asn val gly leu pro gln trp arg cys trp ser pro ala cys 212/71 182/61 TCG GTC ACC TGA CGG TCG GGA TGT TCC TTG GGT CTC GGG TTG CTG GGT TTG CTC AAT ser val thr OPA arg ser gly cys ser leu gly leu gly leu leu gly leu leu asn 272/91 242/81 GCC CTG CTG GTG CGG CGT TCG GCC GAG TCG ATC ACC GCC AAA GAG CAC CCG TTA AAA CGG ala leu leu val arg arg ser ala glu ser ile thr ala lys glu his pro leu lys arg 332/111 302/101 TCG ATG GCC CTC AAC TCG GCA TCG CGA CTG GCG ATT ATC ACC ATG CCT CGG GCT GAT C ser met ala leu asn ser ala ser arg leu ala ile ile thr met pro arg ala asp

SEQ ID No.17B

FIGURE 17B



33/11 GGG CCA CCG ATC AGT CGA TCG GGT GGT TTC CGC TCC ATC AGC CCG GAA TTG AGG TGC CGC gly pro pro ile ser arg ser gly gly phe arg ser ile ser pro glu leu arg cys arg 93/31 63/21 AGT GAC GAC ACC AGC GCA GGA CGC GCC GTT GGT GTT TCC CTC TGT TGC TTT CCC GTC CGG ser asp asp thr ser ala gly arg ala val gly val ser leu cys cys phe pro val arg 153/51 CTC GCC TTT TTT TCA TCA ACG TTG GAC TGC CGC AGT GGC GAT GTT GGT CGC CGG CGT GTT leu ala phe phe ser ser thr leu asp cys arg ser gly asp val gly arg arg arg val 213/71 CGG TCA CCT GAC GGT CGG GAT GTT CCT TGG GTC TCG GGT TGC TGG GTT TGC TCA ATG arg ser pro asp gly arg asp val pro trp val ser gly cys cys trp val cys ser met 273/91 243/81 CCC TGC TGG TGC GGC GTT CGG CCG AGT CGA TCA CCG CCA AAG AGC ACC CGT TAA AAC GGT pro cys trp cys gly val arg pro ser arg ser pro pro lys ser thr arg OCH asn gly 333/111 303/101 CGA TGG CCC TCA ACT CGG CAT CGC GAC TGG CGA TTA TCA CCA TGC CTC GGG CTG ATC arg trp pro ser thr arg his arg asp trp arg leu ser pro cys leu gly leu ile

SEQ ID No.17C

FIGURE 17C

part of the nucleotide sequence of seq17A

31/11 1/1 ggc tag aac ccc gaa gga gac ctc gcg ggt tgc cgg ccc ccg gcc cat cgg atg cgt atc gly AMB asn pro glu gly asp leu ala gly cys arg pro pro ala his arg met arg ile 91/31 61/21 cgg tcg cgc cga ttc acg acc gac ata ggg agc tac ccc ttg ggt gat tcc ggt gcg acg arg ser arg arg phe thr thr asp ile gly ser tyr pro leu gly asp ser gly ala thr 151/51 act gcg ata cgc tcg gcg ggc cac cga tca gtc gat cgg gtg gtt tcc gct cca tca gcc thr ala ile arg ser ala gly his arg ser val asp arg val val ser ala pro ser ala 211/71 cgg aat tga ggt gcc gca gtg acg aca cca gcg cag gac gcg ccg ttg gtg ttt ccc tct arg asn OPA gly ala ala val thr thr pro ala gln asp ala pro leu val phe pro ser 271/91 gtt gct ttc cgt ccg gtt cgc ctt ttt ttc atc aac gtt gga ctg gcc gca gtg gcg atg val ala phe arg pro val arg leu phe phe ile asn val gly leu ala ala val ala met 331/111 301/101 ttg gtc gcc ggc gtg ttc ggt cac ctg acg gtc ggg atg ttc ttg ggt ctc ggg ttg ctg leu val ala gly val phe gly his leu thr val gly met phe leu gly leu gly leu leu 391/131 ctg ggt ttg ctc aat gcc ctg ctg gtg cgg cgt tcg gcc gag tcg atc acc gcc aaa gag leu gly leu leu asn ala leu leu val arg arg ser ala glu ser ile thr ala lys glu 451/151 421/141 cac ccg tta aaa cgg tcg atg gcc ctc aac tcg gca tcg cga ctg gcg att atc acc atc his pro leu lys arg ser met ala leu asn ser ala ser arg leu ala ile ile thr ile 481/161 ctc ggg ctg atc leu gly leu ile

SEQ ID No.17A'

FIGURE 17A'



31/111/1 get aga acc eeg aag gag ace teg egg gtt gee gge eec egg eec ate gga tge gta tee ala arg thr pro lys glu thr ser arg val ala gly pro arg pro ile gly cys val ser 91/31 61/21ggt cgc gcc gat tca cga ccg aca tag gga gct acc cct tgg gtg att ccg gtg cga cga gly arg ala asp ser arg pro thr AMB gly ala thr pro trp val ile pro val arg arg 151/51 121/41 ctg cga tac gct cgg cgg gcc acc gat cag tcg atc ggg tgg ttt ccg ctc cat cag ccc leu arg tyr ala arg arg ala thr asp gln ser ile gly trp phe pro leu his gln pro 211/71 181/61 gga att gag gtg ccg cag tga cga cac cag cgc agg acg cgc cgt tgg tgt ttc cct ctg gly ile glu val pro gln OPA arg his gln arg arg thr arg arg trp cys phe pro leu 271/91 ttg ctt tcc gtc cgg ttc gcc ttt ttt tca tca acg ttg gac tgg ccg cag tgg cga tgt leu leu ser val arg phe ala phe phe ser ser thr leu asp trp pro gln trp arg cys 331/111 301/101 tgg tcg ccg gcg tgt tcg gtc acc tga cgg tcg gga tgt tct tgg gtc tcg ggt tgc tgc trp ser pro ala cys ser val thr OPA arg ser gly cys ser trp val ser gly cys cys 391/131 361/121 tgg gtt tgc tca atg ccc tgc tgg tgc ggc gtt cgg ccg agt cga tca ccg cca aag agc trp val cys ser met pro cys trp cys gly val arg pro ser arg ser pro pro lys ser 451/151 421/141 acc cgt taa aac ggt cga tgg ccc tca act cgg cat cgc gac tgg cga tta tca cca tcc thr arg OCH asn gly arg trp pro ser thr arg his arg asp trp arg leu ser pro ser 481/161 tcg ggc tga tc ser gly OPA

SEQ ID No.17B'

FIGURE 17B'

31/11 1/1 cta gaa ccc cga agg aga cct cgc ggg ttg ccg gcc ccc ggc cca tcg gat gcg tat ccg leu glu pro arg arg arg pro arg gly leu pro ala pro gly pro ser asp ala tyr pro 91/31 61/21 gtc gcg ccg att cac gac cga cat agg gag cta ccc ctt ggg tga ttc cgg tgc gac gac val ala pro ile his asp arg his arg glu leu pro leu gly OPA phe arg cys asp asp 151/51 121/41 tgc gat acg ctc ggc ggg cca ccg atc agt cga tcg ggt ggt ttc cgc tcc atc agc ccg cys asp thr leu gly gly pro pro ile ser arg ser gly gly phe arg ser ile ser pro 211/71 181/61 gaa ttg agg tgc cgc agt gac gac acc agc gca gga cgc gcc gtt ggt gtt tcc ctc tgt glu leu arg cys arg ser asp asp thr ser ala gly arg ala val gly val ser leu cys 271/91 241/81 tgc ttt ccg tcc ggt tcg cct ttt ttt cat caa cgt tgg act ggc cgc agt ggc gat gtt cys phe pro ser gly ser pro phe phe his gln arg trp thr gly arg ser gly asp val 331/111 301/101 ggt cgc cgg cgt gtt cgg tca cct gac ggt cgg gat gtt ctt ggg tct cgg gtt gct gct gly arg arg arg val arg ser pro asp gly arg asp val leu gly ser arg val ala ala 391/131 ggg ttt gct caa tgc cct gct ggt gcg gcg ttc ggc cga gtc gat cac cgc caa aga gca gly phe ala gln cys pro ala gly ala ala phe gly arg val asp his arg gln arg ala 451/151 421/141 ccc gtt aaa acg gtc gat ggc cct caa ctc ggc atc gcg act ggc gat tat cac cat cct pro val lys thr val asp gly pro gln leu gly ile ala thr gly asp tyr his his pro 481/161 cgg gct gat c arg ala asp

SEQ ID No.17C'

FIGURE 17C'

REPLACEMENT SHEET (RULE 26)



```
sequence Rv1303 predicted by Cole et al. (Nature 393:537-544) and partially containing
Seq17A'
                                        31/11
1/1
atg acg aca cca gcg cag gac gcg ccg ttg gtg ttt ccc tct gtt gct ttc cgt ccg gtt
met thr thr pro ala gln asp ala pro leu val phe pro ser val ala phe arg pro val
                                        91/31
61/21
ege ett ttt tte ate aac gtt gga etg gee gea gtg geg atg ttg gte gee gge gtg tte
arg leu phe phe ile asn val gly leu ala ala val ala met leu val ala gly val phe
                                        151/51
121/41
ggt cac ctg acg gtc ggg atg ttc ttg ggt ctc ggg ttg ctg ctg ggt ttg ctc aat gcc
gly his leu thr val gly met phe leu gly leu gly leu leu gly leu leu asn ala
                                        211/71
ctg ctg gtg cgg cgt tcg gcc gag tcg atc acc gcc aaa gag cac ccg tta aaa cgg tcg
leu leu val arg arg ser ala glu ser ile thr ala lys glu his pro leu lys arg ser
                                        271/91
241/81
atg gcc ctc aac tcg gca tcg cga ctg gcg att atc acc atc ctc ggg ctg atc atc gcc
met ala leu asn ser ala ser arg leu ala ile ile thr ile leu gly leu ile ile ala
                                        331/111
tac att ttc cgg ccc gct gga ttg ggc gtc gtg ttc ggg ctg gca ttc ttc cag gtg ctg
tyr ile phe arg pro ala gly leu gly val val phe gly leu ala phe phe gln val leu
                                        391/131
361/121
ctg gtg gca acg acg gcc ctg ccg gtc ctg aag aag ctg cgc act gcg acc gag gaa ccg
leu val ala thr thr ala leu pro val leu lys lys leu arg thr ala thr glu glu pro
                                        451/151
gtc gca act tat tct tcc aat ggc cag acc ggg gga tcg gaa gga agg agc gcc agc gat
val ala thr tyr ser ser asn gly gln thr gly gly ser glu gly arg ser ala ser asp
481/161
gac tga
asp OPA
```

SEQ ID No.17D

FIGURE 17D

```
Orf according to Cole et al. (Nature 393:537-544) and containing Rv1303
                                        31/11
1/1
tga ggt gcc gca gtg acg aca cca gcg cag gac gcg ccg ttg gtg ttt ccc tct gtt gct
OPA gly ala ala val thr thr pro ala gln asp ala pro leu val phe pro ser val ala
                                        91/31
61/21
ttc cgt ccg gtt cgc ctt ttt ttc atc aac gtt gga ctg gcc gca gtg gcg atg ttg gtc
phe arg pro val arg leu phe phe ile asn val gly leu ala ala val ala met leu val
                                        151/51
gcc ggc gtg ttc ggt cac ctg acg gtc ggg atg ttc ttg ggt ctc ggg ttg ctg ctg ggt
ala gly val phe gly his leu thr val gly met phe leu gly leu gly leu leu leu gly
                                        211/71
ttg ctc aat gcc ctg ctg gtg cgg cgt tcg gcc gag tcg atc acc gcc aaa gag cac ccg
leu leu asn ala leu leu val arg arg ser ala glu ser ile thr ala lys glu his pro
                                         271/91
241/81
tta aaa cgg tcg atg gcc ctc aac tcg gca tcg cga ctg gcg att atc acc atc ctc ggg
leu lys arg ser met ala leu asn ser ala ser arg leu ala ile ile thr ile leu gly
                                         331/111
301/101
ctg atc atc gcc tac att ttc cgg ccc gct gga ttg ggc gtc gtg ttc ggg ctg gca ttc
leu ile ile ala tyr ile phe arg pro ala gly leu gly val val phe gly leu ala phe
                                         391/131
361/121
ttc cag gtg ctg ctg gtg gca acg acg gcc ctg ccg gtc ctg aag aag ctg cgc act gcg
phe gln val leu leu val ala thr thr ala leu pro val leu lys lys leu arg thr ala
                                         451/151
421/141
acc gag gaa ccg gtc gca act tat tct tcc aat ggc cag acc ggg gga tcg gaa gga agg
thr glu glu pro val ala thr tyr ser ser asn gly gln thr gly gly ser glu gly arg
481/161
agc gcc agc gat gac tga
ser ala ser asp asp OPA
```

SEQ ID No.17F

FIGURE 17F



31/11 GTC GAA CAG GTA CGG AAG GCG CCG TCG GTC GCT CGG TCC GCT GGT ATC TCG TGT TCA GCC val qlu gln val arg lys ala pro ser val ala arg ser ala gly ile ser cys ser ala 61/21 91/31 AGC CAG CGG CCG TTA ACG TGG CCG AAC AGG TCG TCT TGG GGT CGG GCA TCA GCG TCG ATG ser gln arg pro leu thr trp pro asn arg ser ser trp gly arg ala ser ala ser met 151/51 TGG CTC AGG TCG ATA CCC GAG GGG ATG GCA AGT GTC ACC CCG CCA TCC TTC CAC CTC TTT trp leu arg ser ile pro glu gly met ala ser val thr pro pro ser phe his leu phe 211/71 181/61 ser gly ala thr ile gly pro cys leu thr gly ser arg ala ser his arg pro lys lys 271/91 241/81 ATG CGG AAG ACG ACT CGC GGC CCG ACG CCG CGG AGG CCG CGG CCG AAC CCA AAT CAT met arg lys thr thr arg gly pro thr pro arg arg pro pro arg pro asn pro asn his 301/101 331/111 CAG CCG GTC CCG ATG TTC TCG ACC TAC GGT ATC GCC TCG ACA CTA CTC GGC GTG CTA TCG gln pro val pro met phe ser thr tyr gly ile ala ser thr leu leu gly val leu ser 361/121 GTC GCC GCG GTC GTG CTG GGT GCG ATG ATC val ala ala val val leu gly ala met ile

SEQ ID No.18A

FIGURE 18A

32/11 TCG AAC AGG TAC GGA AGG CGC CGT CGG TCG CTC GGT CCG CTG GTA TCT CGT GTT CAG CCA ser asn arg tyr gly arg arg arg ser leu gly pro leu val ser arg val gln pro 62/21 92/31 GCC AGC GGC CGT TAA CGT GGC CGA ACA GGT CGT CTT GGG GTC GGG CAT CAG CGT CGA TGT ala ser gly arg OCH arg gly arg thr gly arg leu gly val gly his gln arg arg cys 152/51 GGC TCA GGT CGA TAC CCG AGG GGA TGG CAA GTG TCA CCC CGC CAT CCT TCC ACC TCT TTT gly ser gly arg tyr pro arg gly trp gln val ser pro arg his pro ser thr ser phe 182/61 212/71 arg val gln arg ser gly his ala OPA arg gly ala glu pro ala thr gly pro arg arg 272/91 TGC GGA AGA CGA CTC GCG GCC CGA CGC CGC GGA GGC CGC CGC GGC CGA ACC CAA ATC ATC cys gly arg arg leu ala ala arg arg gly gly arg arg gly arg thr gln ile ile 302/101 332/111 AGC CGG TCC CGA TGT TCT CGA CCT ACG GTA TCG CCT CGA CAC TAC TCG GCG TGC TAT CGG ser arg ser arg cys ser arg pro thr val ser pro arg his tyr ser ala cys tyr arg 362/121 TCG CCG CGG TCG TGC TGG GTG CGA TGA TC ser pro arg ser cys trp val arg OPA

SEQ ID No.18B

FIGURE 18B

33/11 3/1 CGA ACA GGT ACG GAA GGC GCC GTC GGT CGC TCG GTC CGC TGG TAT CTC GTG TTC AGC CAG arg thr gly thr glu gly ala val gly arg ser val arg trp tyr leu val phe ser gln 93/31 63/21 CCA GCG GCC GTT AAC GTG GCC GAA CAG GTC GTC TTG GGG TCG GGC ATC AGC GTC GAT GTG pro ala ala val asn val ala glu gln val val leu gly ser gly ile ser val asp val 153/51 123/41 GCT CAG GTC GAT ACC CGA GGG GAT GGC AAG TGT CAC CCC GCC ATC CTT CCA CCT CTT TTC ala gln val asp thr arg gly asp gly lys cys his pro ala ile leu pro pro leu phe 213/71 183/61 gly cys asn asp arg ala met pro asp gly glu gln ser gln pro pro ala gln glu asp 273/91 243/81 GCG GAA GAC GAC TCG CGG CCC GAC GCC GCG GAG GCC GCG GCC GAA CCC AAA TCA TCA ala glu asp asp ser arg pro asp ala ala glu ala ala ala glu pro lys ser ser 333/111 GCC GGT CCC GAT GTT CTC GAC CTA CGG TAT CGC CTC GAC ACT ACT CGG CGT GCT ATC GGT ala gly pro asp val leu asp leu arg tyr arg leu asp thr thr arg arg ala ile gly 363/121 CGC CGC GGT CGT GCT GGG TGC GAT GAT C arg arg gly arg ala gly cys asp asp

SEQ ID No.18C

FIGURE 18C

part of the nucleotide sequence of seq18A 31/11 GAA GGC GCC GTC GGT CGC TCG GTC CGC TGG TAT CTC GTG TTC AGC CAG CCA GCG GCC GTT glu gly ala val gly arg ser val arg trp tyr leu val phe ser gln pro ala ala val 91/31 61/21 AAC GTG GCC GAA CAG GTC GTC TTG GGG TCG GGC ATC AGC GTC GAT GTG GCT CAG GTC GAT asn val ala glu gln val val leu gly ser gly ile ser val asp val ala gln val asp 151/51 121/41 ACC CGA GGG GAT GGC AAG TGT CAC CCC GCC ATC CTT CCA CCT CTT TTC GGG TGC AAC GAT thr arg gly asp gly lys cys his pro ala ile leu pro pro leu phe gly cys asn asp 211/71 181/61 arg ala met pro asp gly glu gln ser gln pro pro ala gln glu asp ala glu asp asp 271/91 241/81 ser arg pro asp ala ala glu ala ala ala ala glu pro lys ser ser ala gly pro met 331/111 301/101 TTC TCG ACC TAC GGT ATC GCC TCG ACA CTA CTC GGC GTG CTA TCG GTC GCC GCG GTC GTG phe ser thr tyr gly ile ala ser thr leu leu gly val leu ser val ala ala val val 361/121 CTG GGT GCG ATG ATC leu gly ala met ile

SEQ ID No.18A'

FIGURE 18A'



31/11 1/1CGG AAG GCG CCG TCG GTC GGT CGG TCC GCT GGT ATC TCG TGT TCA GCC AGC CAG CGG CCG arg lys ala pro ser val ala arg ser ala gly ile ser cys ser ala ser gln arg pro 91/31 61/21 TTA ACG TGG CCG AAC AGG TCG TCT TGG GGT CGG GCA TCA GCG TCG ATG TGG CTC AGG TCG leu thr trp pro asn arg ser ser trp gly arg ala ser ala ser met trp leu arg ser 151/51 121/41 ATA CCC GAG GGG ATG GCA AGT GTC ACC CCG CCA TCC TTC CAC CTC TTT TCG GGT GCA ACG ile pro glu gly met ala ser val thr pro pro ser phe his leu phe ser gly ala thr 211/71 ile gly pro cys leu thr gly ser arg ala ser his arg pro lys lys met arg lys thr 271/91 241/81 thr arg gly pro thr pro arg pro pro arg pro asn pro asn his gln pro val arg 331/111 TGT TCT CGA CCT ACG GTA TCG CCT CGA CAC TAC TCG GCG TGC TAT CGG TCG CCG CGG TCG cys ser arg pro thr val ser pro arg his tyr ser ala cys tyr arg ser pro arg ser 361/121 TGC TGG GTG CGA TGA TC cys trp val arg OPA

SEQ ID No.18B'

FIGURE 18B'

31/11 GGA AGG CGC CGT CGG TCG CTC GGT CCG CTG GTA TCT CGT GTT CAG CCA GCC AGC GGC CGT gly arg arg arg ser leu gly pro leu val ser arg val gln pro ala ser gly arg 91/31 61/21 TAA CGT GGC CGA ACA GGT CGT CTT GGG GTC GGG CAT CAG CGT CGA TGT GGC TCA GGT CGA OCH arg gly arg thr gly arg leu gly val gly his gln arg arg cys gly ser gly arg 151/51 TAC CCG AGG GGA TGG CAA GTG TCA CCC CGC CAT CCT TCC ACC TCT TTT CGG GTG CAA CGA tyr pro arg gly trp gln val ser pro arg his pro ser thr ser phe arg val gln arg 181/61 211/71 ser gly his ala OPA arg gly ala glu pro ala thr gly pro arg arg cys gly arg arg 271/91 241/81 leu ala ala arg arg gly gly arg arg gly arg thr gln ile ile ser arg ser asp 331/111 301/101 GTT CTC GAC CTA CGG TAT CGC CTC GAC ACT ACT CGG CGT GCT ATC GGT CGC CGC GGT CGT val leu asp leu arg tyr arg leu asp thr thr arg arg ala ile gly arg arg gly arg 361/121 GCT GGG TGC GAT GAT C ala gly cys asp asp

SEQ ID No.18C'

FIGURE 18C'



sequence Rv0199 predicted by Cole et al. (Nature 393:537-544) and containing

seq18A' 31/11 1/1 Met pro asp gly glu gln ser gln pro pro ala gln glu asp ala glu asp asp ser arg 61/21 91/31 ccc gac gcc gcg gag gcc gcc gcg gcc gaa ccc aaa tca tca gcc ggt ccg atg ttc tcg pro asp ala ala glu ala ala ala ala glu pro lys ser ser ala gly pro met phe ser 151/51 121/41 acc tac ggt atc gcc tcg aca cta ctc ggc gtg cta tcg gtc gcc gcg gtc gtg ctg ggt thr tyr gly ile ala ser thr leu leu gly val leu ser val ala ala val val leu gly 211/71 181/61 geg atg atc tgg tee gea cae ege gat gae tee gge gag egt ace tae etg ace egg gte

241/81

atg ctg acc gcc gct gaa tgg acg gcc gtg ctg atc aac atg aac gcc gac aac atc gat met leu thr ala ala glu trp thr ala val leu ile asn met asn ala asp asn ile asp 301/101

331/111

gcc agc ctg cag cga ctg cac gac gga acg gtc ggt caa ctc aac acc gac ttc gac gct

ala met ile trp ser ala his arg asp asp ser gly glu arg thr tyr leu thr arg val

gcc agc ctg cag cga ctg cac gac gga acg gtc ggt caa ctc aac acc gac ttc gac gct ala ser leu gln arg leu his asp gly thr val gly gln leu asn thr asp phe asp ala 361/121 391/131

gtc gtg cag ccc tac cgg cag gtg gtg gag aag ttg cgg acg cac agc ggc agg atc val val gln pro tyr arg gln val val glu lys leu arg thr his ser ser gly arg ile 421/141 451/151.

gag gcg gta gcg atc gat acg gtg cac cgc gag ctg gat acc cag tcc ggt gcc gcc cga glu ala val ala ile asp thr val his arg glu leu asp thr gln ser gly ala ala arg 481/161 511/171

ccg gta gta acc acg aaa ttg cca ccg ttt gcc act cgc acc gac tcg gtg ctg ctg gtc pro val val thr thr lys leu pro pro phe ala thr arg thr asp ser val leu leu val 541/181 571/191

gcg acg tcg gtc agt gag aac gcc ggc gcc aaa ccc cag acc gtg cac tgg aac ttg cgg ala thr ser val ser glu asn ala gly ala lys pro gln thr val his trp asn leu arg 601/201 631/211

ctc gat gtc tcc gat gtg gac ggc aag ctg atg atc tcc cgg ttg gag tcg att cga tga leu asp val ser asp val asp gly lys leu met ile ser arg leu glu ser ile arg OPA

SEQ ID No.18D

FIGURE 18D



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ORF according to Cole et al. (Nature 393:537-544) and containing Rv0199
                                        31/11
1/1
taa too gat goo gga ttg ggt gaa atg cao caa gta acg ggt cga gto ttt gga atc ggt
OCH ser asp ala gly leu gly glu met his gln val thr gly arg val phe gly ile gly
                                        91/31
atc gac ata gac tcc gat gcc gcc cac gcc ggc acg ttg cag agt gcc aag ggc ggc
ile asp ile asp ser asp ala ala ala his ala gly thr leu gln ser ala lys gly gly
                                        151/51
121/41
ggc caa ttc ggt ggc gtc ggc cgc gct gtc aat cgt ggc caa ttc gtc gtg cag cgg ttg
gly gln phe gly gly val gly arg ala val asn arg gly gln phe val val gln arg leu
                                        211/71
181/61
cac coe tgc gcg ctc gac ggc ttc ctc gtc gag gaa gct ggc gta gag gtc gcc gat gcg
his pro cys ala leu asp gly phe leu val glu glu ala gly val glu val ala asp ala
                                        271/91
ctg cgc atc ggt gcc tac cgc agc acc tgc ttg gct ggc ctg gat gat cag gtc tcg cac
leu arg ile gly ala tyr arg ser thr cys leu ala gly leu asp asp gln val ser his
                                        331/111
301/101
ttg tgt ctc ggc gcg gtc gaa cag gct acg gaa ggc gcc gtc ggt cgc tcg gtc cgc tgg
leu cys leu gly ala val glu gln ala thr glu gly ala val gly arg ser val arg trp
                                        391/131
361/121
tat ctc gtg ttc agc cag cca gcg gcc gtt aac gtg gcc gaa cag gtc gtc ttg ggg tcg
tyr leu val phe ser gln pro ala ala val asn val ala glu gln val val leu gly ser
                                         451/151
421/141
ggc atc agc gtc gat gtg gct cag gtc gat acc cga ggg gat ggc aag tgt cac ccc gcc .
gly ile ser val asp val ala gln val asp thr arg gly asp gly lys cys his pro ala
                                        511/171
481/161
atc ctt cca cct ctt ttc ggg tgc aac gat cgg gcc atg cct gac ggg gag cag agc cag
ile leu pro pro leu phe gly cys asn asp arg ala met pro asp gly glu gln ser gln
                                        571/191
541/181
cca ccg gcc caa gaa gat gcg gaa gac gac tcg cgg ccc gac gcc gcg gag gcc gcc gcg
pro pro ala gln glu asp ala glu asp asp ser arg pro asp ala ala glu ala ala ala
                                         631/211
601/201
gcc gaa ccc aaa tca tca gcc ggt ccg atg ttc tcg acc tac ggt atc gcc tcg aca cta
ala glu pro lys ser ser ala gly pro met phe ser thr tyr gly ile ala ser thr leu
                                         691/231
661/221
ctc ggc gtg cta tcg gtc gcc gcg gtc gtg ctg ggt gcg atg atc tgg tcc gca cac cgc
leu gly val leu ser val ala ala val val leu gly ala met ile trp ser ala his arg
                                         751/251
gat gac tee gge gag egt ace tae etg ace egg gte atg etg ace gee get gaa tgg acg
asp asp ser gly glu arg thr tyr leu thr arg val met leu thr ala ala glu trp thr
                                         811/271
gcc gtg ctg atc aac atg aac gcc gac aac atc gat gcc agc ctg cag cga ctg cac gac
ala val leu ile asn met asn ala asp asn ile asp ala ser leu gln arg leu his asp
                                         871/291
841/281
gga acg gtc ggt caa ctc aac acc gac ttc gac gct gtc gtg cag ccc tac cgg cag gtg
gly thr val gly gln leu asn thr asp phe asp ala val val gln pro tyr arg gln val
                                         931/311
901/301
gtg gag aag ttg cgg acg cac agc agc ggc agg atc gag gcg gta gcg atc gat acg gtg
val glu lys leu arg thr his ser ser gly arg ile glu ala val ala ile asp thr val
                                         991/331
cac cgc gag ctg gat acc cag tcc ggt gcc gcc cga ccg gta gta acc acg aaa ttg cca
his arg glu leu asp thr gln ser gly ala ala arg pro val val thr thr lys leu pro
                                         1051/351
1021/341
ccg ttt gcc act cgc acc gac tcg gtg ctg ctg gtc gcg acg tcg gtc agt gag aac gcc
pro phe ala thr arg thr asp ser val leu leu val ala thr ser val ser glu asn ala
                                         1111/371
 1081/361
 ggc gcc aaa ccc cag acc gtg cac tgg aac ttg cgg ctc gat gtc tcc gat gtg gac ggc
 gly ala lys pro gln thr val his trp asn leu arg leu asp val ser asp val asp gly
                                         1171/391
 1141/381
 aag ctg atg atc tcc cgg ttg gag tcg att cga tga
 lys leu met ile ser arg leu glu ser ile arg OPA
```

SEQ ID No.18F

FIGURE 18F



31/11 GTT GCG CAA CGG GGT GAG CAC CGA CGC GAT GAT GGC GCA ACT ATC GAA ACT GCA GGA CAT val ala gln arg gly glu his arg arg asp asp gly ala thr ile glu thr ala gly his 91/31 61/21 CGC CAA CGC CAA CGA CGG CAC TCG CGC GGT GGG CAC CCC TGG CTA TCA GGC CAG CGT CGA arg gln arg gln arg arg his ser arg gly gly his pro trp leu ser gly gln arg arg 151/51 CTA TGT GGT AAA CAC ACT GCG CAA CAG CGG TTT TGA TGT GCA AAC CCC GGA GTT CTC CGC leu cys gly lys his thr ala gln gln arg phe OPA cys ala asn pro gly val leu arg 211/71 181/61 TCG CGT GTT CAA GGC CGA AAA AGG GGT GGT GAC CCT CGG CGG CAA CAC CGT GGA GGC GAG ser arg val gln gly arg lys arg gly gly asp pro arg arg gln his arg gly glu 271/91 241/81 GGC GCT CGA GTA CAG CCT CGG CAC ACC GCC GGA CGG GGT GAC GGG CCC GCT GGT GGC TGC gly ala arg val gln pro arg his thr ala gly arg gly asp gly pro ala gly gly cys 331/111 301/101 CCC CGC CGA CGA CAG TCC GGG CTG CAG TCC GTC GGA CTA CGA CAG GCT GCC GGT GTC CGG pro arg arg gln ser gly leu gln ser val gly leu arg gln ala ala gly val arg 361/121 TGC GGT GGT GGT AGA TC cys gly gly ala gly arg

SEQ ID No.19A

FIGURE 19A

32/11 TTG CGC AAC GGG GTG AGC ACC GAC GCG ATG ATG GCG CAA CTA TCG AAA CTG CAG GAC ATC leu arg asn gly val ser thr asp ala met met ala gln leu ser lys leu gln asp ile 62/21 92/31 GCC AAC GCC AAC GAC GGC ACT CGC GCG GTG GGC ACC CCT GGC TAT CAG GCC AGC GTC GAC ala asn ala asn asp gly thr arg ala val gly thr pro gly tyr gln ala ser val asp 152/51 TAT GTG GTA AAC ACA CTG CGC AAC AGC GGT TTT GAT GTG CAA ACC CCG GAG TTC TCC GCT tyr val val asn thr leu arg asn ser gly phe asp val gln thr pro glu phe ser ala 212/71 182/61 CGC GTG TTC AAG GCC GAA AAA GGG GTG GTG ACC CTC GGC GGC AAC ACC GTG GAG GCG AGG arg val phe lys ala glu lys gly val val thr leu gly gly asn thr val glu ala arg 272/91 242/81 GCG CTC GAG TAC AGC CTC GGC ACA CCG CCG GAC GGG GTG ACG GGC CCG CTG GTG GCT GCC ala leu glu tyr ser leu gly thr pro pro asp gly val thr gly pro leu val ala ala 302/101 332/111 CCC GCC GAC GAC AGT CCG GGC TGC AGT CCG TCG GAC TAC GAC AGG CTG CCG GTG TCC GGT pro ala asp asp ser pro gly cys ser pro ser asp tyr asp arg leu pro val ser gly 362/121 GCG GTG GTG CTG GTA GAT C ala val val leu val asp

SEQ ID No.19B

FIGURE 19B



33/11 TGC GCA ACG GGG TGA GCA CCG ACG CGA TGA TGG CGC AAC TAT CGA AAC TGC AGG ACA TCG cys ala thr gly OPA ala pro thr arg OPA trp arg asn tyr arg asn cys arg thr ser 93/31 63/21 CCA ACG CCA ACG ACG GCA CTC GCG CGG TGG GCA CCC CTG GCT ATC AGG CCA GCG TCG ACT pro thr pro thr thr ala leu ala arg trp ala pro leu ala ile arg pro ala ser thr 153/51 ATG TGG TAA ACA CAC TGC GCA ACA GCG GTT TTG ATG TGC AAA CCC CGG AGT TCT CCG CTC met trp OCH thr his cys ala thr ala val leu met cys lys pro arg ser ser pro leu 213/71 183/61 GCG TGT TCA AGG CCG AAA AAG GGG TGG TGA CCC TCG GCG GCA ACA CCG TGG AGG CGA GGG ala cys ser arg pro lys lys gly trp OPA pro ser ala ala thr pro trp arg arg gly 273/91 CGC TCG AGT ACA GCC TCG GCA CAC CGC CGG ACG GGG TGA CGG GCC CGC TGG TGG CTG CCC arg ser ser thr ala ser ala his arg arg thr gly OPA arg ala arg trp trp leu pro 333/111 303/101 CCG CCG ACG ACA GTC CGG GCT GCA GTC CGT CGG ACT ACG ACA GGC TGC CGG TGT CCG GTG pro pro thr thr val arg ala ala val arg arg thr thr thr gly cys arg cys pro val 363/121 CGG TGG TGC TGG TAG ATC arg trp cys trp AMB ile

SEQ ID No.19C

FIGURE 19C

part of the nucleotide sequence of seq19A

31/11 1/1 CTA TCG AAA CTG CAG GAC ATC GCC AAC GCC AAC GGC ACT CGC GCG GTG GGC ACC CCT leu ser lys leu gln asp ile ala asn ala asn asp gly thr arg ala val gly thr pro 91/31 61/21 GGC TAT CAG GCC AGC GTC GAC TAT GTG GTA AAC ACA CTG CGC AAC AGC GGT TTT GAT GTG gly tyr gln ala ser val asp tyr val val asn thr leu arg asn ser gly phe asp val 151/51 CAA ACC CCG GAG TTC TCC GCT CGC GTG TTC AAG GCC GAA AAA GGG GTG GTG ACC CTC GGC gln thr pro glu phe ser ala arg val phe lys ala glu lys gly val val thr leu gly 211/71 GGC AAC ACC GTG GAG GCG AGG GCG CTC GAG TAC AGC CTC GGC ACA CCG CCG GAC GGG GTG gly asn thr val glu ala arg ala leu glu tyr ser leu gly thr pro pro asp gly val 271/91 ACG GGC CCG CTG GTG GCT GCC CCC GCC GAC GAC AGT CCG GGC TGC AGT CCG TCG GAC TAC thr gly pro leu val ala ala pro ala asp asp ser pro gly cys ser pro ser asp tyr 331/111 301/101 GAC AGG CTG CCG GTG TCC GGT GCG GTG GTG CTG GTA GAT C asp arg leu pro val ser gly ala val val leu val asp

SEO ID No.19A'

FIGURE 19A



31/11 1/1 TAT CGA AAC TGC AGG ACA TCG CCA ACG CCA ACG GCA CTC GCG CGG TGG GCA CCC CTG tyr arg asn cys arg thr ser pro thr pro thr thr ala leu ala arg trp ala pro leu 91/31 GCT ATC AGG CCA GCG TCG ACT ATG TGG TAA ACA CAC TGC GCA ACA GCG GTT TTG ATG TGC ala ile arg pro ala ser thr met trp OCH thr his cys ala thr ala val leu met cys 151/51 AAA CCC CGG AGT TCT CCG CTC GCG TGT TCA AGG CCG AAA AAG GGG TGG TGA CCC TCG GCG lys pro arg ser ser pro leu ala cys ser arg pro lys lys gly trp OPA pro ser ala 211/71 181/61 GCA ACA CCG TGG AGG CGA GGG CGC TCG AGT ACA GCC TCG GCA CAC CGC CGG ACG GGG TGA ala thr pro trp arg arg gly arg ser ser thr ala ser ala his arg arg thr gly OPA 241/81 271/91 CGG GCC CGC TGG TGG CTG CCC CCG CCG ACG ACA GTC CGG GCT GCA GTC CGT CGG ACT ACG arg ala arg trp trp leu pro pro pro thr thr val arg ala ala val arg arg thr thr 331/111 301/101 ACA GGC TGC CGG TGT CCG GTG CGG TGG TGC TGG TAG ATC thr gly cys arg cys pro val arg trp cys trp AMB ile

SEQ ID No.19B'

FIGURE 19B'

31/11 1/1 ATC GAA ACT GCA GGA CAT CGC CAA CGC CAA CGG CAC TCG CGC GGT GGG CAC CCC TGG ile glu thr ala gly his arg gln arg gln arg his ser arg gly gly his pro trp 61/21 91/31 CTA TCA GGC CAG CGT CGA CTA TGT GGT AAA CAC ACT GCG CAA CAG CGG TTT TGA TGT GCA leu ser gly gln arg arg leu cys gly lys his thr ala gln gln arg phe OPA cys ala 151/51 121/41 AAC CCC GGA GTT CTC CGC TCG CGT GTT CAA GGC CGA AAA AGG GGT GGT GAC CCT CGG CGG asn pro gly val leu arg ser arg val gln gly arg lys arg gly gly asp pro arg arg 211/71 181/61 CAA CAC CGT GGA GGC GAG GGC GCT CGA GTA CAG CCT CGG CAC ACC GCC GGA CGG GGT GAC gln his arg gly gly glu gly ala arg val gln pro arg his thr ala gly arg gly asp 271/91 GGG CCC GCT GGT GGC TGC CCC CGC CGA CGA CAG TCC GGG CTG CAG TCC GTC GGA CTA CGA gly pro ala gly gly cys pro arg arg gln ser gly leu gln ser val gly leu arg 331/111 301/101 CAG GCT GCC GGT GTC CGG TGC GGT GGT GGT AGA TC gln ala ala gly val arg cys gly gly ala gly arg

SEQ ID No.19C'

FIGURE 19C'



sequence Rv0418 predicted by Cole et al. (Nature 393:537-544) and containing seq19A'

1/1 31/11atg gtg aac aaa too agg atg atg cog gcg gtg ctg gcc gtg gct gtg gtc gtc gca ttc Met val asn lys ser arg met met pro ala val leu ala val ala val val ala phe 91/31 ctg acg acg ggc tgt atc cgg tgg tct acg cag tcg cgg ccc gtt gtt aac ggc ccc gct leu thr thr gly cys ile arg trp ser thr gln ser arg pro val val asn gly pro ala 151/51 121/41 gcc gca gag ttc gcc gtt gcg ttg cgc aac cgg gtg agc acc gac gcg atg atg gcg cac ala ala glu phe ala val ala leu arg asn arg val ser thr asp ala met met ala his 211/71 181/61 cta tcg aaa ctg cag gac atc gcc aac gcc aac gac ggc act cgc gcg gtg ggc acc cct leu ser lys leu gln asp ile ala asn ala asn asp gly thr arg ala val gly thr pro 271/91 241/81 ggc tat cag gcc agc gtc gac tat gtg gta aac aca ctg cgc aac agc ggt ttt gat gtg gly tyr gln ala ser val asp tyr val val asn thr leu arg asn ser gly phe asp val 331/111 301/101 caa acc ccg gag ttc tcc gct cgc gtg ttc aag gcc gaa aaa ggg gtg gtg acc ctc ggc gln thr pro glu phe ser ala arg val phe lys ala glu lys gly val val thr leu gly 391/131 ggc aac acc gtg gag gcg agg gcg ctc gag tac agc ctc ggc aca ccg ccg gac ggg gtg gly asn thr val glu ala arg ala leu glu tyr ser leu gly thr pro pro asp gly val 421/141 451/151 acg ggc ccg ctg gtg gct gcc ccc gcc gac gac agt ccg ggc tgc agt ccg tcg gac tac thr gly pro leu val ala ala pro ala asp asp ser pro gly cys ser pro ser asp tyr 511/171 qac agg ctg ccg gtg tcc ggt gcg gtg gtg ctg gta gat cgc ggc gtc tgt cct ttt gcc asp arg leu pro val ser gly ala val leu val asp arg gly val cys pro phe ala 571/191 541/181 cag aag gaa gac gca gcc gcg cag cgc ggt gcg gtg gcg ctg atc att gct gac aac atc gln lys glu asp ala ala ala gln arg gly ala val ala leu ile ile ala asp asm ile 631/211 601/201 gac gag cag gcg atg ggc ggc acc ctg ggg gct aat acc gac gtc aag atc ccg gtg gtg asp glu gln ala met gly gly thr leu gly ala asn thr asp val lys ile pro val val 691/231 661/221 agt gtc acc aag tcg gtc gga ttc cag cta cgc gga cag tct ggg cca acc acc gtc aag ser val thr lys ser val gly phe gln leu arg gly gln ser gly pro thr thr val lys 751/251 721/241 ctc acg gcg agc acc caa agt ttc aag gcc cgc aac gtc atc gcg cag acg aag acg ggg leu thr ala ser thr gln ser phe lys ala arg asn val ile ala gln thr lys thr gly 811/271 781/261 tcg tcg gcc aac gtg gtg atg gca ggt gcg cat ttg gac agc gtt ccg gaa gga ccc ggc ser ser ala asn val val met ala gly ala his leu asp ser val pro glu gly pro gly 871/291 841/281 atc aac gac aac ggc tcg gga gtg gct gcg gtt ctg gaa acg gca gtg cag ctg ggg aac ile asn asp asn gly ser gly val ala ala val leu glu thr ala val gln leu gly asn 931/311 tca ccg cat gtg tcc aac gcg gta cgg ttc gcc ttc tgg ggc gcc gag gaa ttc ggc ctg ser pro his val ser asn ala val arg phe ala phe trp gly ala glu glu phe gly leu 991/331 961/321 att ggg tca cga aac tac gtc gag tcg ctg gac atc gac gcg ctc aaa ggc atc gcg ctg ile gly ser arg asn tyr val glu ser leu asp ile asp ala leu lys gly ile ala leu

SEQ ID No.19 D

FIGURE 19D

ORF according to Cole et al. (Nature 393:537-544) and containing Rv0418

```
31/11
1/1
tag gcc att caa cgc tct gtt cgt ttg att ggt cgg tgg gat gcg aaa gct gcg cga
AMB ala ile gln arg ser val arg leu ile gly arg trp asp ala lys ala ala arg arg
cag gcg cgg tct aat ctg ggc gcg atg gtg aac aaa tcc agg atg atg ccg gcg gtg ctg
gln ala arg ser asn leu gly ala met val asn lys ser arg met met pro ala val leu
                                        151/51
gcc gtg gct gtg gtc gtc gca ttc ctg acg acg ggc tgt atc cgg tgg tct acg cag tcg
ala val ala val val val ala phe leu thr thr gly cys ile arg trp ser thr gln ser
                                        211/71
cgg ccc gtt gtt aac ggc ccc gct gcc gca gag ttc gcc gtt gcg ttg cgc aac cgg gtg
arg pro val val asn gly pro ala ala ala glu phe ala val ala leu arg asn arg val
                                        271/91
241/81
ago aco gao gog atg atg gog cao ota tog aaa otg cag gao ato goo aac goo aac gao
ser thr asp ala met met ala his leu ser lys leu gln asp ile ala asn ala asn asp
                                        331/111
ggc act cgc gcg gtg ggc acc cct ggc tat cag gcc agc gtc gac tat gtg gta aac aca
gly thr arg ala val gly thr pro gly tyr gln ala ser val asp tyr val val asn thr
                                        391/131
361/121
ctg cgc aac agc ggt ttt gat gtg caa acc ccg gag ttc tcc gct cgc gtg ttc aag gcc
leu arg asn ser gly phe asp val gln thr pro glu phe ser ala arg val phe lys ala
                                        451/151
421/141
gaa aaa ggg gtg gtg acc ctc ggc ggc aac acc gtg gag gcg agg gcg ctc gag tac agc
glu lys gly val val thr leu gly gly asn thr val glu ala arg ala leu glu tyr ser
                                        511/171
481/161
ctc ggc aca ccg ccg gac ggg gtg acg ggc ccg ctg gtg gct gcc ccc gcc gac gac agt
leu gly thr pro pro asp gly val thr gly pro leu val ala ala pro ala asp asp ser
                                        571/191
ccg ggc tgc agt ccg tcg gac tac gac agg ctg ccg gtg tcc ggt gcg gtg gtg ctg gta
pro gly cys ser pro ser asp tyr asp arg leu pro val ser gly ala val val leu val
                                        631/211
gat cgc ggc gtc tgt cct ttt gcc cag aag gaa gac gca gcc gcg cag cgc ggt gcg gtg
asp arg gly val cys pro phe ala gln lys glu asp ala ala ala gln arg gly ala val
                                         691/231
661/221
gcg ctg atc att gct gac aac atc gac gag cag gcg atg ggc ggc acc ctg ggg gct aat
ala leu ile ile ala asp asn ile asp glu gln ala met gly gly thr leu gly ala asn
                                         751/251
acc gac gtc aag atc ccg gtg gtg agt gtc acc aag tcg gtc gga ttc cag cta cgc gga
thr asp val lys ile pro val val ser val thr lys ser val gly phe gln leu arg gly
                                         811/271
781/261
cag tot ggg coa acc acc gto aag cto acg gcg ago acc caa agt tto aag gco cgc aac
gln ser gly pro thr thr val lys leu thr ala ser thr gln ser phe lys ala arg asn
                                         871/291
gtc atc gcg cag acg aag acg ggg tcg tcg gcc aac gtg gtg atg gca ggt gcg cat ttg
val ile ala gln thr lys thr gly ser ser ala asn val val met ala gly ala his leu
                                         931/311
901/301
gac ago gtt cog gaa gga coo ggo ato aac gac aac ggo tog gga gtg got gog gtt otg
asp ser val pro glu gly pro gly ile asn asp asn gly ser gly val ala ala val leu
                                         991/331
gaa acg gca gtg cag ctg ggg aac tca ccg cat gtg tcc aac gcg gta cgg ttc gcc ttc
glu thr ala val gln leu gly asn ser pro his val ser asn ala val arg phe ala phe
```

SEQ ID No.19 F

FIGURE 19F

4



1051/351 1021/341 tgg ggc gcc gag gaa ttc ggc ctg att ggg tca cga aac tac gtc gag tcg ctg gac atc trp gly ala glu glu phe gly leu ile gly ser arg asn tyr val glu ser leu asp ile 1111/371 1081/361 gac gcg ctc aaa ggc atc gcg ctg tat ctg aac ttc gac atg ttg gcg tcg ccg aac ccg asp ala leu lys gly ile ala leu tyr leu asn phe asp met leu ala ser pro asn pro 1171/391 1141/381 ggt tac ttc acc tac gac ggt gac cag tcg ctg ccg cta gac gcc cgc ggt cag ccg gtg gly tyr phe thr tyr asp gly asp gln ser leu pro leu asp ala arg gly gln pro val 1231/411 1201/401 gtg ccc gaa ggc tcg gcc ggt atc gag cgc acg ttc gtc gcc tat ctg aag atg gcc ggc val pro glu gly ser ala gly ile glu arg thr phe val ala tyr leu lys met ala gly 1291/431 1261/421 aag acc gcg cag gac acc tcg ttc gac ggt cgg tcc gac tac gac ggc ttc acg ctg gcg lys thr ala gln asp thr ser phe asp gly arg ser asp tyr asp gly phe thr leu ala 1351/451 1321/441 ggt atc cct tcg ggt ggc ctg ttc tcc ggc gct gag gtc aag aag tcc gcc gag caa gcc gly ile pro ser gly gly leu phe ser gly ala glu val lys lys ser ala glu gln ala 1411/471 gag ctc tgg ggc ggc acc gcc gac gag cct ttc gat ccc aac tat cac cag aag aca gac glu leu trp gly gly thr ala asp glu pro phe asp pro asn tyr his gln lys thr asp 1471/491 1441/481 acc ctg gac cat atc gac cgc acc gcg ctc ggt atc aac ggc gct ggc gtc gcg tac gcg thr leu asp his ile asp arg thr ala leu gly ile asn gly ala gly val ala tyr ala 1531/511 gtg ggt ttg tat gcg cag gac ctc ggc ggc ccc aac ggg gtt ccg gtc atg gcg gac cgc val gly leu tyr ala gln asp leu gly gly pro asn gly val pro val met ala asp arg 1561/521 acc cgc cac ctg att gcc aaa ccg tga thr arg his leu ile ala lys pro OPA

SEQ ID No.19F (continued)

FIGURE 19F (continued)

31/11 CGA GAC AGT GGT GCG GGA CAC TTG AGT TCG GCT GCT AAC GAC GCC AGA GTC GCC CGC TTC arg asp ser gly ala gly his leu ser ser ala ala asn asp ala arg val ala arg phe 91/31 61/21 CGC GGT GTG GGA CTC ACG TTC GGT GAG GGT ACA GCG GAC CTT CGA GCA CGC AAT ATC GTG arg gly val gly leu thr phe gly glu gly thr ala asp leu arg ala arg asn ile val 151/51 121/41 GGC CGG CTG GCA ACC GTC GGT TTC GAC GTT GGT GAC GAC CCC TCG TTC ATG AAT CGT TCT gly arg leu ala thr val gly phe asp val gly asp asp pro ser phe met asn arg ser 211/71 181/61 TGA GCT CCC CGT TTT GCT GGA TGC CCA GGC ACC GCC GGT ACT GCT GCG CTT AAG CTT GTC OPA ala pro arg phe ala gly cys pro gly thr ala gly thr ala ala leu lys leu val 271/91 241/81 GCA CAT GGT GCC GGC AGG GAA CAG TGG GCA AGC AGC TAG CCG CGC TCG CCG CGC TGG ala his gly ala gly arg glu glu gln trp ala ser ser AMB pro arg ser pro arg trp 331/111 301/101 TCG GTG CGT GCA TGC TCG CAG CCG GAT GCA CCA ACG TGG TCG ACG GGA CCG CCG TGG CTG ser val arg ala cys ser gln pro asp ala pro thr trp ser thr gly pro pro trp leu 361/121 CCG ACA AAT CCG GAC CAC TGC ATC AGG ATC pro thr asn pro asp his cys ile arg ile

SEQ ID No.20A

FIGURE 20A
REPLACEMENT SHEET (RULE 26)

32/11

GAG ACA GTG GTG CGG GAC ACT TGA GTT CGG CTG CTA ACG ACG CCA GAG TCG CCC GCT TCC qlu thr val val arg asp thr OPA val arg leu leu thr thr pro glu ser pro ala ser 92/31 62/21 GCG GTG TGG GAC TCA CGT TCG GTG AGG GTA CAG CGG ACC TTC GAG CAC GCA ATA TCG TGG ala val trp asp ser arg ser val arg val gln arg thr phe glu his ala ile ser trp 152/51 122/41 GCC GGC TGG CAA CCG TCG GTT TCG ACG TTG GTG ACG ACC CCT CGT TCA TGA ATC GTT CTT ala gly trp gln pro ser val ser thr leu val thr thr pro arg ser OPA ile val leu 212/71 GAG CTC CCC GTT TTG CTG GAT GCC CAG GCA CCG CCG GTA CTG CTG CGC TTA AGC TTG TCG glu leu pro val leu leu asp ala gln ala pro pro val leu leu arg leu ser leu ser 272/91 242/81 CAC ATG GTG CCG GCA GGG AGG AAC AGT GGG CAA GCA GCT AGC CGC GCT CGC CGC GCT GGT his met val pro ala gly arg asn ser gly gln ala ala ser arg ala arg ala gly 332/111 CGG TGC GTG CAT GCT CGC AGC CGG ATG CAC CAA CGT GGT CGA CGG GAC CGC CGT GGC TGC arg cys val his ala arg ser arg met his gln arg gly arg arg asp arg gly cys 362/121 CGA CAA ATC CGG ACC ACT GCA TCA GGA TC arg gln ile arg thr thr ala ser gly

SEQ ID No.20B

FIGURE 20B

33/11

AGA CAG TGG TGC GGG ACA CTT GAG TTC GGC TGC TAA CGA CGC CAG AGT CGC CCG CTT CCG arg gln trp cys gly thr leu glu phe gly cys OCH arg arg gln ser arg pro leu pro 93/31 63/21 CGG TGT GGG ACT CAC GTT CGG TGA GGG TAC AGC GGA CCT TCG AGC ACG CAA TAT CGT GGG arg cys gly thr his val arg OPA gly tyr ser gly pro ser ser thr gln tyr arg gly 153/51 123/41 CCG GCT GGC AAC CGT CGG TTT CGA CGT TGG TGA CGA CCC CTC GTT CAT GAA TCG TTC TTG pro ala gly asn arg arg phe arg arg trp OPA arg pro leu val his glu ser phe leu 183/61 213/71 AGC TCC CCG TTT TGC TGG ATG CCC AGG CAC CGC CGG TAC TGC TGC GCT TAA GCT TGT CGC ser ser pro phe cys trp met pro arg his arg arg tyr cys cys ala OCH ala cys arg 273/91 243/81 ACA TGG TGC CGG CAG GGA ACA GTG GGC AAG CAG CTA GCC GCG CTC GCC GCG CTG GTC thr trp cys arg gln gly gly thr val gly lys gln leu ala ala leu ala ala leu val 333/111 GGT GCG TGC ATG CTC GCA GCC GGA TGC ACC AAC GTG GTC GAC GGG ACC GCC GTG GCT gly ala cys met leu ala ala gly cys thr asn val val asp gly thr ala val ala ala 363/121 GAC AAA TCC GGA CCA CTG CAT CAG GAT C asp lys ser gly pro leu his gln asp

SEQ ID No.20C

FIGURE 20C



part of the nucleotide sequence of seq20A

1/1									31/1	11								
TGT GGG	ACT	CAC	GTT	CGG	TGA	GGG	TAC	AGC	GGA	CCT	TCG	AGC	ACG	CAA	TAT	CGT	GGG	CCG
cys gly	thr	his	val	arg	OPA	gly	tyr	ser	gly	pro	ser	ser	thr	gln	tyr	arg	gly	pro
61/21									91/3	31								
GCT GGC	AAC	CGT	CGG	TTT	CGA	CGT	TGG	TGA	CGA	CCC	CTC	GTT	CAT	GAA	TCG	TTC	TTG	AGC
ala gly	asn	arg	arg	phe	arg	arg	trp	OPA	arg	pro	leu	val	his	glu	ser	phe	leu	ser
121/41									151/	51								
TCC CCG	TTT	TGC	TGG	ATG	CCC	AGG	CAC	CGC	CGG	TAC	TGC	TGC	GCT	TAA	GCT	TGT	CGC	ACA
ser pro	phe	cys	trp	met	pro	arg	his	arg	arg	tyr	cys	cys	ala	OCH	ala	cys	arg	thr
181/61									211/	71								
TGG TGC																		
trp cys	arg	gln	gly	gly	thr	val	gly	lys	gln	leu	ala	ala	leu	ala	ala	leu	val	gly
241/81									271/									
GCG TGC																		
ala cys	met	leu	ala	ala	gly	cys	thr	asn	val	val	asp	gly	thr	ala	val	ala	ala	asp
301/101																		
AAA TCC	GGA	CCA	CTG	CAT	CAG	GAT	С											
lys ser	gly	pro	leu	his	gln	asp												

SEQ ID No.20A'

FIGURE 20A'

1/1									31/1	11								
GTG GGA	CTC	ACG	TTC	GGT	GAG	GGT	ACA	GCG	GAC	CTT	CGA	GCA	CGC	AAT	ATC	GTG	GGC	CGG
val gly	leu	thr	phe	gly	glu	gly	thr	ala	asp	leu	arg	ala	arg	asn	ile	val	gly	arg
61/21									91/3	31								
CTG GCA	ACC	GTC	GGT	TTC	GAC	GTT	GGT	GAC	GAC	CCC	TCG	TTC	ATG	AAT	CGT	TCT	TGA	GCT
leu ala	thr	val	gly	phe	asp	val	gly	asp	asp	pro	ser	phe	met	asn	arg	ser	OPA	ala
121/41									151,	/51								
CCC CGT	-																	
pro arg	phe	ala	gly	cys	pro	gly	thr	ala	gly	thr	ala	ala	leu	lys	leu	val	ala	his
181/61									211,	. —								
GGT GCC																		
gly ala	gly	arg	glu	glu	gln	trp	ala	ser	ser	AMB	pro	arg	ser	pro	arg	trp	ser	val
241/81									271,	/91								
CGT GCA	TGC	TCG	CAG	CCG	GAT	GCA	CCA	ACG	TGG	TCG	ACG	GGA	CCG	CCG	TGG	CTG	CCG	ACA
arg ala	cys	ser	gln	pro	asp	ala	pro	thr	trp	ser	thr	gly	pro	pro	trp	leu	pro	thr
301/101																		
AAT CCG	GAC	CAC	TGC	ATC	AGG	ATC												
asn pro	asp	his	cys	ile	arg	ile												

SEQ ID No.20B'

FIGURE 20B'



31/11 1/1 GTG TGG GAC TCA CGT TCG GTG AGG GTA CAG CGG ACC TTC GAG CAC GCA ATA TCG TGG GCC val trp asp ser arg ser val arg val gln arg thr phe glu his ala 1le ser trp ala 91/31 GGC TGG CAA CCG TCG GTT TCG ACG TTG GTG ACG ACC CCT CGT TCA TGA ATC GTT CTT GAG gly trp gln pro ser val ser thr leu val thr thr pro arg ser OPA ile val leu glu 151/51 121/41 CTC CCC GTT TTG CTG GAT GCC CAG GCA CCG CCG GTA CTG CTG CGC TTA AGC TTG TCG CAC leu pro val leu leu asp ala gln ala pro pro val leu leu arg leu ser leu ser his 211/71 ATG GTG CCG GCA GGG AGG AAC AGT GGG CAA GCA GCT AGC CGC GCT CGC CGC GCT CGG met val pro ala gly arg asn ser gly gln ala ala ser arg ala arg arg ala gly arg 271/91 TGC GTG CAT GCT CGC AGC CGG ATG CAC CAA CGT GGT CGA CGG GAC CGC CGT GGC TGC CGA cys val his ala arg ser arg met his gln arg gly arg arg asp arg gly cys arg 301/101 CAA ATC CGG ACC ACT GCA TCA GGA TC gln ile arg thr thr ala ser gly

SEQ ID No.20C'

FIGURE 20C'

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sequence Rv3576 predicted by Cole et al. (Nature 393:537-544) and containing seq20A'
                                        31/11
1/1
atg ggc aag cag cta gcc gcg ctc gcc gcg ctg gtc ggt gcg tgc atg ctc gca gcc gga
met gly lys gln leu ala ala leu ala ala leu val gly ala cys met leu ala ala gly
                                        91/31
tgc acc aac gtg gtc gac ggg acc gcc gtg gct gcc gac aaa tcc gga cca ctg cat cag
cys thr asn val val asp gly thr ala val ala ala asp lys ser gly pro leu his gln
                                        151/51
gat ccg ata ccg gtt tca gcg ctt gaa ggg ctg ctt ctc gac ttg agc cag atc aat gcc
asp pro ile pro val ser ala leu glu gly leu leu leu asp leu ser gln ile asn ala
                                        211/71
gcg ctg ggt gcg aca tcg atg aag gtg tgg ttc aac gcc aag gca atg tgg gac tgg agc
ala leu gly ala thr ser met lys val trp phe asn ala lys ala met trp asp trp ser
                                        271/91
241/81
aag agc gtg gcc gac aag aat tgc ctg gct atc gac ggt cca gca cag gaa aag gtc tat
lys ser val ala asp lys asn cys leu ala ile asp gly pro ala gln glu lys val tyr
                                        331/111
301/101
gcc ggc acc ggg tgg acc gct atg cgc ggc caa cgg ctg gat gac agc atc gat gac tcc
ala gly thr gly trp thr ala met arg gly gln arg leu asp asp ser ile asp asp ser
                                         391/131
361/121
aag aaa cgc gac cac tac gcc att caa gcg gtc gtc ggc ttc ccg acc gca cat gat gcc
lys lys arg asp his tyr ala ile gln ala val val gly phe pro thr ala his asp ala
                                         451/151
421/141
gag gag ttc tac agc tcc tcg gtg caa agc tgg agc agc tgc tcg aac cgc cgg ttt gtc
glu glu phe tyr ser ser ser val gln ser trp ser ser cys ser asn arg arg phe val
                                         511/171
481/161
gaa gtc acc ccc gga cag gac gac gcc gcc tgg act gtg gct gac gtt gtc aac gac aac
glu val thr pro gly gln asp asp ala ala trp thr val ala asp val val asn asp asn
                                         571/191
541/181
ggc atg ctc agt agc tcg cag gtt cag gaa ggc ggc gac gga tgg acc tgc cag cgt gcc
gly met leu ser ser ser gln val gln glu gly gly asp gly trp thr cys gln arg ala
                                         631/211
ctg act gcg cgc aac aac gtc act atc gac att gtc acg tgc gcc tat agc caa ccg gat
leu thr ala arg asn asn val thr ile asp ile val thr cys ala tyr ser gln pro asp
                                         691/231
661/221
ttg gtg gcg att ggc atc gct aac caa atc gcg gcc aag gtt gct aag cag tag
leu val ala ile gly ile ala asn gln ile ala ala lys val ala lys gln AMB
```

SEQ ID No.20D

FIGURE 20D



ORF according to Cole et al. (Nature 393:537-544) and containing Rv3576

1/1									31/1									
taa gct	tgt	cgc	aca	tgg	tgc	cgg	cag	gga	gga	aca	gtg	ggc	aag	cag	cta	gcc	gcg	ctc
OCH ala	cys	arg	thr	trp	cys	arg	gln	gly			val	gly	lys	gln	leu	aia	ala	leu
61/21					.		a+ a	~	91/3	_	+~~	200	220	~+ <i>~</i>	a+a	~~~	~~~	200
gcc gcg ala ala	ctg	gtc	ggt	gcg	cys	met	1 - 11	gCa ala	ala	gga	CVS	thr	asn	val	val	asp	alv	thr
121/41	reu	Val	g r y	ата	Cys	nicc	1.Cu	ulu	151/		CID	CIII	abii	• • • •	V 4.1	авр	9-1	0111
gcc gtg	act	acc	gac	aaa	tcc	qqa	cca	ctg			gat	ccg	ata	ccg	gtt	tca	gcg	ctt
ala val	ala	ala	asp	lys	ser	gly	pro	leu	his	gln	asp	pro	ile	pro	val	ser	ala	leu
181/61									211/	71								
gaa ggg	ctg	ctt	ctc	gac	ttg	agc	cag	atc	aat	gcc	gcg	ctg	ggt	gcg	aca	tcg	atg	aag
glu gly	leu	leu	leu	asp	leu	ser	gln	ile			ala	leu	gly	ala	thr	ser	met	lys
241/81							+ ~ ~	~~~	271/		226	200	~+ ~	aca	a 2 a	224		taa
gtg tgg val trp	ttc	aac	gcc	aag	gca	atg met	trn	gac	trn	agc	lve	ear	y c y	ala	asn	lve	aat	CVS
301/101	pne	asii	ата	тур	ата	mec	стр	asp	331/		TYS	501	vui	ala	авр	T Y S	asii	CYS
ctg gct	atc	gac	aat	cca	σca	caq	gaa	aaq			qcc	aac	acc	aga	tgg	acc	gct	atg
leu ala	ile	asp	gly	pro	ala	gln	glu	lys	val	tyr	āla	gly	thr	gly	trp	thr	āla	met
361/121									391,	131							-	
cgc ggc	caa	cgg	ctg	gat	gac	agc	atc	gat	gac	tcc	aag	aaa	cgc	gac	cac	tac	gcc	att
arg gly	gln	arg	leu	asp	asp	ser	ile	asp			lys	lys	arg	asp	his	tyr	ala	ile
421/141										151					. ~ .	+ ~ ~	+~~	~+~
caa gcg gln ala	gtc	gtc	ggc	ttc	ccg	acc	gca	cat	gat	gcc	gag	gag	nhe	tac	agc	cer	ser	g cg val
gin aia 481/161	Val	Val	дту	pne	pro	CILL	ara	1112		171	gru	gru	pire	CAT	SCL	501	501	vai
caa agc	taa	agc	agc	tac	t.ca	aac	cac	caa	-		gaa	atc	acc	ccc	qqa	caq	gac	qac
gln ser	trp	ser	ser	cys	ser	asn	arg	arg	phe	val	glu	val	thr	pro	gly	gln	asp	asp
541/181									571,	/191								
gcc gcc	tgg	act	gtg	gct	gac	gtt	gtc	aac	gac	aac	ggc	atg	ctc	agt	agc	tcg	cag	gtt
ala ala	trp	thr	val	ala	asp	val	val	asn			gly	met	leu	ser	ser	ser	gln	val
601/201										/211								
cag gaa	ggc	ggc	gac	gga	tgg	acc	tgc	cag	cgt	gcc	ctg	act +hr	gcg	cgc	aac	aac	gtc	thr
gln glu 661/221	дтА	gry	asp	gry	crb	CILL	Cys	9111		/231	100	CIII	ara	arg	asn	abii	Vul	CILL
atc gac	att	atc	aca	tac	acc	tat	agg	caa			tta	ata	aca	att	aac	atc	act	aac
ile asp	ile	val	thr	CVS	ala	tyr	ser	gln	pro	asp	leu	val	ala	ile	gly	ile	āla	asn
721/241			-	.		-		_	-	-					- -			
caa atc																		
gln ile	ala	ala	lys	val	ala	lys	gln	AMB										

SEQ ID No.20F

FIGURE 20F



1/1							31/3	11								
GTC CTG GTC	GCC GC	G CAA	CTG	GCC	GGT	CCC	GAT	GGA	AAG	TGT	TCA	CGA	TCG	CGC	TTC	TGC
val leu val	ala al	a gln	leu	ala	gly	pro	asp	gly	lys	cys	ser	arg	ser	arg	phe	cys
61/21							91/3	31								
CGC TGG TAG	TGG CG	A TGG	TGT	TAG	CAG	GAT	TGC	GGG	TCG	AGG	CTG	CGA	TGG	CCA	GCA	CCA
arg trp AMB	trp ar	g trp	cys	AMB	gln	asp	cys	gly	ser	arg	leu	arg	trp	pro	ala	pro
121/41							151,	/51								
GCG GCC TGC	GGC TG	G TCG	CCG	CGC	GCG	CCG	AAA	TGA	TAC	CCG	CGA	TCA	CGA	AAT	ACA	TGT
ala ala cys	gly tr	p ser	pro	arg	ala	pro	_		tyr	pro	arg	ser	arg	asn	thr	cys
181/61							211,	/71								
CGG CGC TGG	ACG TO	G CCG	TGC	TGG	CCA	GCT	CGA	CCG	GAC	ACG	ATG	TGG	AGG	GGG	CGC	AGA
arg arg trp	thr se	r pro	cys	trp	pro	ala	arg	pro	asp	thr	met	trp	arg	gly	arg	arg
241/81							271,	/91								
AAA ACT TCA	CCG CC	C GCA	AGT	ACG	AGC	TGC	AGA	CGC	GAC	TGG	CCG	ACA	CCG	ACG	TCA	TCG
lys thr ser	pro pr	o ala	ser	thr	ser	cys	arg	arg	asp	trp	pro	thr	pro	thr	ser	ser
301/101							331,	/111								
CAG ACG TGC																
gln thr cys	gly ar	g glu	OPA	thr	arg	cys	ser	thr	ala	val	arg	arg	cys	trp	ile	arg
361/121																
TGC TGG CCG	ACA GO	A TCG	GCT	TGC	GGG	ATC										
cys trp pro	thr al	a ser	ala	cys	gly	ile									-	

SEQ ID No.21A

FIGURE 21A

32/11 TCC TGG TCG CCG CGC AAC TGG CCG GTC CCG ATG GAA AGT GTT CAC GAT CGC GCT TCT GCC ser trp ser pro arg asn trp pro val pro met glu ser val his asp arg ala ser ala 92/31 62/21 GCT GGT AGT GGC GAT GGT GTT AGC AGG ATT GCG GGT CGA GGC TGC GAT GGC CAG CAC CAG ala gly ser gly asp gly val ser arg ile ala gly arg gly cys asp gly gln his gln 152/51 CGG CCT GCG GCT GGT CGC CGC GCG CGC CGA AAT GAT ACC CGC GAT CAC GAA ATA CAT GTC arq pro ala ala gly arg arg ala arg arg asn asp thr arg asp his glu ile his val 182/61 212/71 GGC GCT GGA CGT CGC CGT GCT GGC CAG CTC GAC CGG ACA CGA TGT GGA GGG GGC GCA GAA gly ala gly arg arg ala gly gln leu asp arg thr arg cys gly gly ala glu 272/91 AAA CTT CAC CGC CCG CAA GTA CGA GCT GCA GAC GCG ACT GGC CGA CAC CGA CGT CAT CGC lys leu his arg pro gln val arg ala ala asp ala thr gly arg his arg arg his arg 332/111 AGA CGT GCG GTC GGG AGT GAA CAC GCT GCT CAA CGG CGG TCA GGC GCT GCT GGA TAA GAT arg arg ala val gly ser glu his ala ala gln arg arg ser gly ala ala gly OCH asp 362/121 GCT GGC CGA CAG CAT CGG CTT GCG GGA TC ala gly arg gln his arg leu ala gly

SEQ ID No.21B

FIGURE 21B





33/11 CCT GGT CGC CGC GCA ACT GGC CGG TCC CGA TGG AAA GTG TTC ACG ATC GCG CTT CTG CCG pro gly arg arg ala thr gly arg ser arg trp lys val phe thr ile ala leu leu pro 63/21 93/31 CTG GTA GTG GCG ATG GTG TTA GCA GGA TTG CGG GTC GAG GCT GCG ATG GCC AGC AGC leu val val ala met val leu ala gly leu arg val glu ala ala met ala ser thr ser 153/51 GGC CTG CGG CTG GTC GCC GCG CGC GCC GAA ATG ATA CCC GCG ATC ACG AAA TAC ATG TCG gly leu arg leu val ala ala arg ala glu met ile pro ala ile thr lys tyr met ser 183/61 213/71 GCG CTG GAC GTC GCC GTG CTG GCC AGC TCG ACC GGA CAC GAT GTG GAG GGG GCG CAG AAA ala leu asp val ala val leu ala ser ser thr gly his asp val glu gly ala gln lys 243/81 273/91 AAC TTC ACC GCC CGC AAG TAC GAG CTG CAG ACG CGA CTG GCC GAC ACC GAC GTC ATC GCA asn phe thr ala arg lys tyr glu leu gln thr arg leu ala asp thr asp val ile ala 303/101 333/111 GAC GTG CGG TCG GGA GTG AAC ACG CTG CTC AAC GGC GGT CAG GCG CTG CTG GAT AAG ATG asp val arg ser gly val asn thr leu leu asn gly gly gln ala leu leu asp lys met 363/121 CTG GCC GAC AGC ATC GGC TTG CGG GAT C

SEQ ID No.21C

FIGURE 21C

part of the nucleotide sequence of seq21A

leu ala asp ser ile gly leu arg asp

1/1 31/11 ACG ATC GCG CTT CTG CCG CTG GTA GTG GCG ATG GTG TTA GCA GGA TTG CGG GTC GAG GCT thr ile ala leu leu pro leu val val ala met val leu ala gly leu arg val glu ala 91/31 GCG ATG GCC AGC ACC AGC GGC CTG CGG CTG GTC GCC GCG CGC GAA ATG ATA CCC GCG ala met ala ser thr ser gly leu arg leu val ala ala arg ala glu met ile pro ala 121/41 151/51 ATC ACG AAA TAC ATG TCG GCG CTG GAC GTC GCC GTG CTG GCC AGC TCG ACC GGA CAC GAT ile thr lys tyr met ser ala leu asp val ala val leu ala ser ser thr gly his asp 211/71 GTG GAG GGG GCG CAG AAA AAC TTC ACC GCC CGC AAG TAC GAG CTG CAG ACG CGA CTG GCC val glu gly ala gln lys asn phe thr ala arg lys tyr glu leu gln thr arg leu ala 241/81 271/91 GAC ACC GAC GTC ATC GCA GAC GTG CGG TCG GGA GTG AAC ACG CTG CTC AAC GGC GGT CAG asp thr asp val ile ala asp val arg ser gly val asn thr leu leu asn gly gly gln 301/101 331/111 GCG CTG CTG GAT AAG ATG CTG GCC GAC AGC ATC GGC TTG CGG GAT C ala leu leu asp lys met leu ala asp ser ile gly leu arg asp

SEQ ID No.21A'

FIGURE 21A'





sequence Rv3365c predicted by Cole et al. (Nature 393:537-544) and containing Seq21A'

31/11 1/1 gtg acc atg ttc gcc cgc ccg acc atc ccg gtc gcg gcc gct tct gat att tcc gcc val thr met phe ala arg pro thr ile pro val ala ala ala ala ser asp ile ser ala 91/31 ccg gct caa ccg gcc cgc ggc aaa cct cag caa cgc ccg ccg tcc tgg tcg ccg cgc aac pro ala gln pro ala arg gly lys pro gln gln arg pro pro ser trp ser pro arg asn 151/51 121/41 tgg ccg gtc cga tgg aaa gtg ttc acg atc gcg ctt ctg ccg ctg gta gtg gcg atg gtg trp pro val arg trp lys val phe thr ile ala leu leu pro leu val val ala met val 211/71 tta gca gga ttg cgg gtc gag gct gcg atg gcc agc acc agc ggc ctg cgg ctg gtc qcc leu ala gly leu arg val glu ala ala met ala ser thr ser gly leu arg leu val ala 271/91 geg ege gee gaa atg ata eee geg ate aeg aaa tae atg teg geg etg gae gte gee gtg ala arg ala glu met ile pro ala ile thr lys tyr met ser ala leu asp val ala val 331/111 301/101 ctg gcc agc tcg acc gga cac gat gtg gag ggg gcg cag aaa aac ttc acc gcc cgc aag leu ala ser ser thr gly his asp val glu gly ala gln lys asn phe thr ala arg lys 391/131 tac gag ctg cag acg cga ctg gcc gac acc gac gtc atc gca gac gtg cgg tcg gga gtg tyr glu leu gln thr arg leu ala asp thr asp val ile ala asp val arg ser gly val 451/151 421/141 aac acg ctg ctc aac ggc ggt cag gcg ctg ctg gat aag gtg ctg gcc gac agc atc ggc asn thr leu leu asn gly gly gln ala leu leu asp lys val leu ala asp ser ile gly 511/171 481/161 ttg cgg gat cgg gtc acc gcc tac gcg ccg ctg ctg ttg acg gcc cag aac gtg att gac leu arg asp arg val thr ala tyr ala pro leu leu leu thr ala gln asn val ile asp 571/191 541/181 gcg tcg gtg cgg gtt gac agc gag caa atc cga acc cag gtg cag ggt ttg agc cga gcc ala ser val arg val asp ser glu gln ile arg thr gln val gln gly leu ser arg ala 631/211 601/201 gtt ggc gcc cgc ggg cag atg acg atg cag gag atc ctg gtg act cgc ggc gcc gac ctt val gly ala arg gly gln met thr met gln glu ile leu val thr arg gly ala asp leu 691/231 661/221 gee gag eeg caa etg ege age geg atg gtt ace etg gee gge ace gaa eee teg aeg etg ala glu pro gln leu arg ser ala met val thr leu ala gly thr glu pro ser thr leu 751/251 ttc ggg atg agc gcg gcg ctc ggt gca ggc tcg ccg gac acc aag aac ctg cag caa phe gly met ser ala ala leu gly ala gly ser pro asp thr lys asn leu gln gln 811/271 atg gtg acc agg atg gcg atc atg tcc gat ccg gcc gtt gca ctg gtc aac aac cca gag met val thr arg met ala ile met ser asp pro ala val ala leu val asn asn pro glu 871/291 ctg ctg cac tcg ata cag atc acc cgc gac att gcc gag cag gtg atc acc gac acc leu leu his ser ile gln ile thr arg asp ile ala glu gln val ile thr asp thr thr 931/311 901/301 gag gcg gtg acg aag tcg gtg caa agc cag gcc acc gac cgg cgg gat gcc gcg att cgc glu ala val thr lys ser val gln ser gln ala thr asp arg arg asp ala ala ile arg 991/331 gac gcc gtg ctg gtg ttg gcc gcc atc gcg acc gcg atc gtc gtc gtg ttg gtg gcg asp ala val leu val leu ala ala ile ala thr ala ile val val leu val val ala

SEQ ID No.21F

FIGURE 21D



1861/621								1891									
atc aga cta	cgc	ggt	ccg	gtg	acc	ggt	gaa	cag	ggc	acc	ggc	acc	acc	gcc	gag	gtc	tac
val gly leu	arg	gly	pro	val	thr	gly	glu	gln	gly	thr	gly	thr	thr	ala	glu	val	tyr
1921/641									/651								
ctg ccg cta	gcc	gtg	ctc	gag	ggg	acg	gcc	cca	gcg	cag	ccg	cca	aag	ccg	cgg	gta	ttt
leu pro leu	ala	val	leu	glu	gly	thr	aia				pro	pro	туs	pro	arg	vaı	pne
1981/661					~~~	~~~	~~~		/671		cca	2 C C	aac.	att	ccc	acc	acc
gcg atc aag ala ile lys	ccg	ccg	tgt	222	gaa	nro	gcg	gcg	ala	asn	nro	thr	asp	val	pro	ala	ala
2041/681	bro	pro	Cys	pro	gru	pro	ата	2071	./691	. app	PLO	0111	чор	• • • •	PLU	u_u	414
atc ggg ccg	cta	cca	cca	ata	acq	tta	ctc				acc	ccq	aaa	tcc	agt	ggc	atc
ile gly pro	leu	pro	pro	val	thr	leu	leu	pro	arg	arg	thr	pro	gly	ser	ser	gly	ile
2101/701								2131	./711	L							
acc gac gtc	ccg	gcc	cag	ccg	atg	cag	cag	cgg	cgg	cgc	gag	ctg	aaa	aca	CCC	tgg	tgg
ala asp val	pro	ala	gln	pro	met	gln	gln	arg	arg	arg	glu	leu	lys	thr	pro	trp	trp
2161/721								2191	./731	L							
gag gat agg	ttt	caa	cag	gag	CCC	aaa	caa	ccg	ccc	gca	cca	gaa	ccg	cga	ccg	gcg	ccg
glu asp arg	phe	gln	gln	glu	pro	lys	gln	pro	pro	a⊥a '	pro	glu	pro	arg	pro	ата	pro
2221/741									L/753		~~~	~~~	a+ a	ata	t a.c	200	caa
ccg ccc gcc pro pro ala	aaa	CCC	gcg	cca	ccg	gcg	ggc	nro	gcc	gat	gac	gac	y cc	ile	tur	ara	ard
	Lys	pro	ата	pro	bro	ата	дту	2311	vai 1/77:	asp 1	asp	asp	Val	110	CYL	arg	arg
2281/761 atg ctc tcc	asa	ata	ata	aat	gac	cca	cac				cac	agc	ccc	gat	ctq	gac	tgg
met leu ser	alu	met	val	alv	asp	pro	his	glu	leu	ála	his	ser	pro	asp	leu	asp	trp
2341/781	gra	11100		9-1	<u>F</u> -	L		237	1/79	1			-				
aad tod gtg	taa	gac	cac	ggc	tgg	tcg	gcg	gcc	gcc	gag	gcc	gcg	gac	aag	ccc	gtg	cag
lys ser val	trp	asp	his	gly	trp	ser	ala	ala	ala	glu	ala	ala	asp	lys	pro	val	gln
2401/801								243	1/81:	1							
tcc cgc acg	gac	tac	ggc	ctg	ccg	gtg	cgc	gaa	ccc	ggg	gcc	cgg	tta	gtg	ccg	ggg	gcg
ser arg thr	asp	tyr	gly	leu	pro	val	arg	glu	pro	g⊥y	ala	arg	leu	val	pro	дтА	ата
2461/821									1/83		~~~	at a	~~~	+ ~ ~	220	aac	aaa
gcg gtg cct ala val pro	gag	gga	ccc	gat	cgg	gag	bie	nro	ggu	gca	gcg	len	ala	ser	asn	alv	alv
	gru	g∓Ā	Þτο	asp	ary	gru	111.5	255	1/85	1	ulu		aza			5-1	5-1
2521/841 ctt cat ccc	aac	cas	aca	cca	caa	cac	aca				cac	gac	ccc	gac	qcq	gtt	cgt
leu his pro	al v	ara	ala	pro	ard	his	ala	ala	ala	val	arq	asp	pro	asp	ala	val	arg
2581/861	9-1	4-9		F	5			261	1/87	1	_	_	_	_			
acc tcc atc	aqc	agc	cat	ttc	ggc	ggc	gtg	cgc	acc	ggg	cgg	tcg	cat	gcc	cgc	gag	agc
ala ser ile	ser	ser	his	phe	gly	gly	val	arg	thr	gly	arg	ser	his	ala	arg	glu	ser
2641/881																	
agt cag gga	ccc	aat	cag	caa	tga												
ser gln gly	pro	asn	gln	gln	OPA												

SEQ ID No.21F (continued 2)

FIGURE 21F (continued 2)

31/11 CTA CGA CAA GGC AAA GGA GCA CAG GGT GAA GCG TGG ACT GAC GGT CGC GGT AGC CGG AGC leu arg gln gly lys gly ala gln gly glu ala trp thr asp gly arg gly ser arg ser 91/31 61/21 CGC CAT TCT GGT CGC AGG TCT TTC CGG ATG TTC AAG CAA CAA GTC GAC TAC AGG AAG CGG arg his ser gly arg arg ser phe arg met phe lys gln gln val asp tyr arg lys arg 151/51 121/41 TGA GAC CAC GAC CGC AGG CAC GAC GGC AAG CCC CGG CGC ATC CGG GCC GAA GGT OPA asp his asp arg gly arg his asp gly lys pro arg arg arg ile arg ala glu gly 211/71 181/61 CGT CAT CGA CGG TAA GGA CCA GAA CGT CAC CGG GTC TGT GGT GTG CAC AAC CGC GGC CGG arg his arg arg OCH gly pro glu arg his arg val cys gly val his asn arg gly arg 271/91 CAA TGT CAA CAT CGC GAT CGG CGG GGC GGC CGG CAT TGC CGC CGT GCT CAC CGA CGG gln cys gln his arg asp arg arg gly gly asp arg his cys arg arg ala his arg arg 331/111 301/101 CAA CCC TCC GGA GGT GAA GTC CGT TGG GCT CGG TAA CGT CAA CGG CGT CAC GCT GGG ATA gln pro ser gly gly glu val arg trp ala arg OCH arg gln arg arg his ala gly ile 391/131 361/121 CAC GTC GGG CAC CGG ACA GGG TAA CGC TCG GCA ACC AAG GAC GGC AGC CAC TAC AAG ATC his val gly his arg thr gly OCH arg ser ala thr lys asp gly ser his tyr lys ile

SEQ ID No.22A

FIGURE 22A

32/11 TAC GAC AAG GCA AAG GAG CAC AGG GTG AAG CGT GGA CTG ACG GTC GCG GTA GCC GGA GCC tyr asp lys ala lys glu his arg val lys arg gly leu thr val ala val ala gly ala 92/31 62/21 GCC ATT CTG GTC GCA GGT CTT TCC GGA TGT TCA AGC AAC AAG TCG ACT ACA GGA AGC GGT ala ile leu val ala gly leu ser gly cys ser ser asn lys ser thr thr gly ser gly 152/51 122/41 glu thr thr thr ala ala gly thr thr ala ser pro gly ala ala ser gly pro lys val 212/71 182/61 GTC ATC GAC GGT AAG GAC CAG AAC GTC ACC GGG TCT GTG GTG TGC ACA ACC GCG GCC val ile asp gly lys asp gln asn val thr gly ser val val cys thr thr ala ala gly 272/91 242/81 AAT GTC AAC ATC GCG ATC GGC GGG GCG ACC GGC ATT GCC GCC GTG CTC ACC GAC GGC asn val asn ile ala ile gly gly ala ala thr gly ile ala ala val leu thr asp gly 332/111 302/101 AAC CCT CCG GAG GTG AAG TCC GTT GGG CTC GGT AAC GTC AAC GGC GTC ACG CTG GGA TAC asn pro pro glu val lys ser val gly leu gly asn val asn gly val thr leu gly tyr 392/131 362/121 ACG TCG GGC ACC GGA CAG GGT AAC GCT CGG CAA CCA AGG ACG GCA GCC ACT ACA AGA TC thr ser gly thr gly gln gly asn ala arg gln pro arg thr ala ala thr thr arg

SEQ ID No.22B

FIGURE 22B



33/11 ACG ACA AGG CAA AGG AGC ACA GGG TGA AGC GTG GAC TGA CGG TCG CGG TAG CCG GAG CCG thr thr arg gln arg ser thr gly OPA ser val asp OPA arg ser arg AMB pro glu pro 93/31 63/21 CCA TTC TGG TCG CAG GTC TTT CCG GAT GTT CAA GCA ACA AGT CGA CTA CAG GAA GCG GTG pro phe trp ser gln val phe pro asp val gln ala thr ser arg leu gln glu ala val 153/51 123/41 arg pro arg pro arg gln ala arg arg gln ala pro ala pro his pro gly arg arg ser 213/71 TCA TCG ACG GTA AGG ACC AGA ACG TCA CCG GGT CTG TGG TGT GCA CAA CCG CGG CCG ser ser thr val arg thr arg thr ser pro gly leu trp cys ala gln pro arg pro ala 273/91 243/81 ATG TCA ACA TCG CGA TCG GCG GGG CGG CGA CCG GCA TTG CCG CCG TGC TCA CCG ACG GCA met ser thr ser arg ser ala gly arg arg pro ala leu pro pro cys ser pro thr ala 333/111 ACC CTC CGG AGG TGA AGT CCG TTG GGC TCG GTA ACG TCA ACG GCG TCA CGC TGG GAT ACA thr leu arg arg OPA ser pro leu gly ser val thr ser thr ala ser arg trp asp thr 393/131 363/121 CGT CGG GCA CCG GAC AGG GTA ACG CTC GGC AAC CAA GGA CGG CAG CCA CTA CAA GAT C arg arg ala pro asp arg val thr leu gly asn gln gly arg gln pro leu gln asp

SEQ ID No.22C

FIGURE 22C

31/11 GCA CAA CCG CGG CCG GCA ATG TCA ACA TCG CGA TCG GCG GGG CGA CCG GCA TTG CCG ala gln pro arg pro ala met ser thr ser arg ser ala gly arg arg pro ala leu pro 91/31 CCG TGC TCA CCG ACG GCA ACC CTC CGG AGG TGA AGT CCG TTG GGC TCG GTA ACG TCA ACG pro cys ser pro thr ala thr leu arg arg OPA ser pro leu gly ser val thr ser thr 151/51 121/41 GCG TCA CGC TGG GAT ACA CGT CGG GCA CCG GAC AGG GTA ACG CCT CGG CAA CCA AGG ACG ala ser arg trp asp thr arg arg ala pro asp arg val thr pro arg gln pro arg thr 211/71 GCA GCC ACT ACA AGA TCA CAG GGT GAA GCG TGG ACT GAC GGT CGC GGT AGC CGC ala ala thr thr arg ser gln gly glu ala trp thr asp gly arg gly ser arg ser arg 271/91 241/81 CAT TCT GGT CGC AGG TCT TTC CGG ATG TTC AAG CAA CAA GTC GAC TAC AGG AAG CGG TGA his ser gly arg arg ser phe arg met phe lys gln gln val asp tyr arg lys arg OPA 331/111 301/101 asp his asp arg gly arg his asp gly lys pro arg arg arg ser gly pro lys val val 391/131 361/121 ATC GAC GGT AAG GAC CAG AAC GTC ACC GGC TCC GTG GTG TGC ACA ACC GCG GCC GGC AAT ile asp gly lys asp gln asn val thr gly ser val val cys thr thr ala ala gly asn 451/151 GTC AAC ATC GCG ATC GGC GGG GCG GCG ACC GGC ATT GCC GCC GTG CTC ACC GAC GGC AAC val asn ile ala ile gly gly ala ala thr gly ile ala ala val leu thr asp gly asn 511/171 481/161 CCT CCG GAG GTG AAG TCC GTT GGG CTC GGT AAC GTC AAC GGC GTC ACG CTG GGA TAC ACG pro pro glu val lys ser val gly leu gly asn val asn gly val thr leu gly tyr thr 571/191 541/181 TCG GGC ACC GGA CAG GGT AAC GCC TCG GCA ACC AAG GAC GGC AGC CAC TAC AAG ATC ser gly thr gly gln gly asn ala ser ala thr lys asp gly ser his tyr lys ile

SEQ ID No.23A

FIGURE 23A REPLACEMENT SHEET (RULE 26)





									32/1									
CAC AAC	CGC	GGC	CGG	CAA	TGT	CAA	CAT	CGC	GAT	CGG	CGG	GGC	GGC	GAC	CGG	CAT	TGC	CGC
his asn	arq	qly	arq	gln	cys	gln	his	arg	asp	arg	arg	gly	gly	asp	arg	his	cys	arg
62/21									92/3	31								
CGT GCT	CAC	CGA	CGG	CAA	CCC	TCC	GGA	GGT	GAA	GTC	CGT	TGG	GCT	CGG	TAA	CGT	CAA	CGG
arg ala	his	arg	arg	gln	pro	ser	gly	gly	glu	val	arg	trp	ala	arg	OCH	arg	gln	arg
122/41									152/			~~~	am a	a aa	77.7	~ 7 7	CCA	ccc
CGT CAC	GCT	GGG	ATA	CAC	GTC	GGG	CAC	CGG	ACA	GGG	TAA	CGC	CTC	GGC	AAC	CAA	GGA	250
arg his	ala	gly	ile	his	val	gly	his	arg	thr	g _T y	ОСН	arg	Ieu	дту	asn	gin	дту	arg
182/61						am.a	n n C	ccm	212/		א כי כי	CTC	ccc	CT A	GCC	GGA	GCC	GCC
CAG CCA	CTA	CAA	GAT	CAC	AGG	GTG	AAG	CGT	GGA ~1	101	+hr	1721	212	ual	ala	alv	ala	ala
gln pro	leu	gln	asp	his	arg	vaı	туs	arg	919 272	16u /01	CILL	var	ата	var	ara	9-1	ara	uzu
242/81 ATT CTG	~~~	~~3	C CIT	cmm.	maa	CCA	m/m	ሞሮሽ			AAG	TCG	АСТ	ACA	GGA	AGC	GGT	GAG
ATT CTG	GTC	GCA	GGT	CTT	TCC	GGA	101	CAT	cer	agn	lus	ser	thr	thr	alv	ser	alv	alu
	vaı	aıa	дту	reu	SCI	gry	Cys	301	332	/111	-10				5-2		, 1	,
302/101 ACC ACG	300	ccc	CCA	CCC	ACG	»CG	GCA	AGC			GCC	GCT	CCG	GGC	CGA	AGG	TCG	TCA
thr thr	ACC +hr	212	ala	41 tz	thr	thr	ala	ser	pro	alv	ala	ala	pro	qly	arq	arg	ser	ser
262/121									392,	/131								
መሮር እሮር	СΤΆ	AGG	ACC	AGA	ACG	TCA	CCG	GCT	CCG	TGG	TGT	GCA	CAA	CCG	CGG	CCG	GCA	ATG
ser thr	val	arg	thr	arq	thr	ser	pro	ala	pro	trp	cys	ala	gln	pro	arg	pro	ala	met
122/141									452	/151								
שכא אכא	TCG	CGA	TCG	GCG	GGG	CGG	CGA	CCG	GCA	TTG	CCG	CCG	TGC	TCA	CCG	ACG	GCA	ACC
ser thr	ser	arq	ser	ala	gly	arg	arg	pro	ala	leu	pro	pro	cys	ser	pro	thr	ala	thr
102/161									512	/171								
CTC CGG	AGG	TGA	AGT	CCG	TTG	GGC	TCG	GTA	ACG	TCA	ACG	GCG	TCA	ÇGC	TGG	GAT	ACA	CGT
leu arg	arg	OPA	ser	pro	leu	gly	ser	val	thr	ser	thr	ala	ser	arg	trp	asp	tnr	arg
542/181										/191	7.00	C C 7	ccc	х ст	א כי א	λCλ	T)C	
CGG GCA	CCG	GAC	AGG	GTA	ACG	CCT	CGG	CAA	. CCA	AGG	ACG	GCA	313	+h~	ACA +h~	. AGA	. 10	
arg ala	pro	asp	arg	val	thr	pro	arg	g⊥n	pro	arg	LIII	ата	ата	CILL	LIIL	ary		

SEQ ID No.23B

FIGURE 23B



33/11 ACA ACC GCG GCC GGC AAT GTC AAC ATC GCG ATC GGC GCG GCG ACC GGC ATT GCC GCC thr thr ala ala gly asn val asn ile ala ile gly gly ala ala thr gly ile ala ala 93/31 63/21 GTG CTC ACC GAC GGC AAC CCT CCG GAG GTG AAG TCC GTT GGG CTC GGT AAC GTC AAC GGC val leu thr asp gly asn pro pro glu val lys ser val gly leu gly asn val asn gly 153/51 123/41 GTC ACG CTG GGA TAC ACG TCG GGC ACC GGA CAG GGT AAC GCC TCG GCA ACC AAG GAC GGC val thr leu gly tyr thr ser gly thr gly gln gly asn ala ser ala thr lys asp gly 213/71 183/61 AGC CAC TAC AAG ATC ACA GGG TGA AGC GTG GAC TGA CGG TCG CGG TAG CCG CAG CCA ser his tyr lys ile thr gly OPA ser val asp OPA arg ser arg AMB pro glu pro pro 273/91 TTC TGG TCG CAG GTC TTT CCG GAT GTT CAA GCA ACA AGT CGA CTA CAG GAA GCG GTG AGA phe trp ser gln val phe pro asp val gln ala thr ser arg leu gln glu ala val arg 333/111 303/101 CCA CGA CCG CGG CAG GCA CGA CGG CAA GCC CCG GCG CCC CTC CGG GCC GAA GGT CGT CAT pro arg pro arg gln ala arg arg gln ala pro ala pro leu arg ala glu gly arg his 393/131 CGA CGG TAA GGA CCA GAA CGT CAC CGG CTC CGT GGT GTG CAC AAC CGC GGC CGG CAA TGT arg arg OCH gly pro glu arg his arg leu arg gly val his asn arg gly arg gln cys 453/151 423/141 CAA CAT CGC GAT CGG CGG GGC GGC CGG CAT TGC CGC CGT GCT CAC CGA CGG CAA CCC gln his arg asp arg gly gly asp arg his cys arg arg ala his arg arg gln pro 513/171 483/161 TCC GGA GGT GAA GTC CGT TGG GCT CGG TAA CGT CAA CGG CGT CAC GCT GGG ATA CAC GTC ser gly gly glu val arg trp ala arg OCH arg gln arg arg his ala gly ile his val 573/191 543/181 GGG CAC CGG ACA GGG TAA CGC CTC GGC AAC CAA GGA CGG CAG CCA CTA CAA GAT C gly his arg thr gly OCH arg leu gly asn gln gly arg gln pro leu gln asp

SEQ ID No.23C

FIGURE 23C





31/11 CTA ACG ACA GGC AAA GGA GCA CAG GGT GAA GCG TGG ACT GAC GGT CGC GGT AGC CGG AGC leu thr thr gly lys gly ala gln gly glu ala trp thr asp gly arg gly ser arg ser 91/31 61/21 CGC CAT TCT GGT CGC AGG TCT TTC CGG ATG TTC AAG CAA CAA GTC GAC TAC AGG AAG CGG arg his ser gly arg arg ser phe arg met phe lys gln gln val asp tyr arg lys arg 151/51 OPA asp his asp arg gly arg his asp gly lys pro arg arg arg ser gly pro lys val 211/71 GTC ATC GAC GGT AAG GAC CAG AAC GTC ACC GGC TCC GTG GTG TGC ACA ACC GCG GCC GGC val ile asp gly lys asp gln asn val thr gly ser val val cys thr thr ala ala gly 271/91 AAT GTC AAC ATC GCG ATC GGC GGG GCG ACC GGC ATT GCC GCC GTG CTC ACC GAC GGC asn val asn ile ala ile gly gly ala ala thr gly ile ala ala val leu thr asp gly 331/111 301/101 AAC CCT CCG GAG GTG AAG TCC GTT GGG CTC GGT AAC GTC AAC GGC GTC ACG CTG GGA TAC asn pro pro glu val lys ser val gly leu gly asn val asn gly val thr leu gly tyr 391/131 361/121 ACG TCG GGC ACC GGA CAG GGT AAC GCC TCG GCA ACC AAG GAC GGC AGC CAC TAC AAG ATC thr ser gly thr gly gln gly asn ala ser ala thr lys asp gly ser his tyr lys ile

SEQ ID No.24A

FIGURE 24A

32/11 TAA CGA CAG GCA AAG GAG CAC AGG GTG AAG CGT GGA CTG ACG GTC GCG GTA GCC GGA GCC OCH arg gln ala lys glu his arg val lys arg gly leu thr val ala val ala gly ala 92/31 62/21 GCC ATT CTG GTC GCA GGT CTT TCC GGA TGT TCA AGC AAC AAG TCG ACT ACA GGA AGC GGT ala ile leu val ala gly leu ser gly cys ser ser asn lys ser thr thr qly ser gly 152/51 122/41 glu thr thr thr ala ala gly thr thr ala ser pro gly ala ala pro gly arg arg ser 212/71 182/61 TCA TCG ACG GTA AGG ACC AGA ACG TCA CCG GCT CCG TGG TGT GCA CAA CCG CGG CCG GCA ser ser thr val arg thr arg thr ser pro ala pro trp cys ala gln pro arg pro ala 272/91 242/81 ATG TCA ACA TCG CGA TCG GCG GGG CGG CGA CCG GCA TTG CCG CCG TGC TCA CCG ACG GCA met ser thr ser arg ser ala gly arg arg pro ala leu pro pro cys ser pro thr ala 332/111 302/101 ACC CTC CGG AGG TGA AGT CCG TTG GGC TCG GTA ACG TCA ACG GCG TCA CGC TGG GAT ACA thr leu arg arg OPA ser pro leu gly ser val thr ser thr ala ser arg trp asp thr 392/131 362/121 CGT CGG GCA CCG GAC AGG GTA ACG CCT CGG CAA CCA AGG ACG GCA GCC ACT ACA AGA TC arg arg ala pro asp arg val thr pro arg gln pro arg thr ala ala thr thr arg

SEQ ID No.24B

FIGURE 24B



AAC GAC AGG CAA AGG AGC ACA GGG TGA AGC GTG GAC TGA CGG TCG CGG TAG CCG GAG CCG asn asp arg gln arg ser thr gly OPA ser val asp OPA arg ser arg AMB pro glu pro 93/31 CCA TTC TGG TCG CAG GTC TTT CCG GAT GTT CAA GCA ACA AGT CGA CTA CAG GAA GCG GTG pro phe trp ser gln val phe pro asp val gln ala thr ser arg leu gln glu ala val 153/51 123/41 AGA CCA CGA CCG CGG CAG GCA CGA CGG CAA GCC CCG GCG CCC CTC CGG GCC GAA GGT CGT arg pro arg pro arg gln ala arg arg gln ala pro ala pro leu arg ala glu gly arg 213/71 183/61 CAT CGA CGG TAA GGA CCA GAA CGT CAC CGG CTC CGT GGT GTG CAC AAC CGC GGC CGA his arg arg OCH gly pro glu arg his arg leu arg gly val his asn arg gly arg gln 273/91 243/81 TGT CAA CAT CGC GAT CGG CGG GGC GGC CGC CGT TGC CGC CGT GCT CAC CGA CGG CAA cys gln his arg asp arg gly qly asp arg his cys arg arg ala his arg arg gln 333/111 303/101 CCC TCC GGA GGT GAA GTC CGT TGG GCT CGG TAA CGT CAA CGG CGT CAC GCT GGG ATA CAC pro ser gly gly glu val arg trp ala arg OCH arg gln arg arg his ala gly ile his 393/131 363/121 GTC GGG CAC CGG ACA GGG TAA CGC CTC GGC AAC CAA GGA CGG CAG CCA CTA CAA GAT C val gly his arg thr gly OCH arg leu gly asn gln gly arg gln pro leu gln asp

SEQ ID No.24C

FIGURE 24C

Direct primer

5' ACG CGG CGC AGC CTG TTG 3'

SEQ ID No.25

FIGURE 25

Reverse primer

5' CGA CCT TGG GAT TCG CCT 3'

SEQ ID No.26

FIGURE 26





31/11

CCT ACC AGC AAG AGC CCA GGG CTT CAC AGG ACC TAA AAG GAG TAG CGC CCA TGG GCT TGA pro thr ser lys ser pro gly leu his arg thr OCH lys glu AMB arg pro trp ala OPA 91/31 TCC AAT TTT CCT TCC GCC CCG TGC AAT ACC ATC TGC AAG ACC AGC GAC GGC CCG TGG TTG ser asn phe pro ser ala pro cys asn thr ile cys lys thr ser asp gly pro trp leu 151/51 121/41 CGG TCG CGC AGC TTG CGG AAA CGG GGT ATG GAC CCT GCC GTA CCG TTG TTG CCA CTT GAT arg ser arg ser leu arg lys arg gly met asp pro ala val pro leu leu pro leu asp 211/71 GTC GTC GCT CTC CAC CCG TCG GGG GGC GAA AGC CAT TCC GAC ACT GGG ATC CTC AAA ACG val val ala leu his pro ser gly gly glu ser his ser asp thr gly ile leu lys thr 271/91 TCG GCT GAG TGT CTG CAG GGC TCC GGG GAG CAG CCG ATC ATC ACC ATG TAC GAA CTG AAT ser ala glu cys leu gln gly ser gly glu gln pro ile ile thr met tyr glu leu asn 331/111 AAG TCC CCC CCG CGC GAC TTC CAG ACA TTT GTT GTG GTT TCG GTT GAG GCC GAG GCG AGG lys ser pro pro arg asp phe gln thr phe val val ser val glu ala glu ala arg 391/131 361/121 CTC ATT TCG CAG CAA GCG GTC TCC GGG TCG CAG CAT CGT TGC GGC GAT CGC GGC GCA GTC leu ile ser gln gln ala val ser gly ser gln his arg cys gly asp arg gly ala val 421/141 GTC GGA CGA GTC GTC GTC AAC GAC CAC GAT C val gly arg val val val asn asp his asp

SEQ ID No.27A

FIGURE 27A

31/11

CTA CCA GCA AGA GCC CAG GGC TTC ACA GGA CCT AAA AGG AGT AGC GCC CAT GGG CTT GAT leu pro ala arg ala gln gly phe thr gly pro lys arg ser ser ala his gly leu asp 91/31 61/21 CCA ATT TTC CTT CCG CCC CGT GCA ATA CCA TCT GCA AGA CCA GCG ACG GCC CGT GGT TGC pro ile phe leu pro pro arg ala ile pro ser ala arg pro ala thr ala arg gly cys 151/51 121/41 GGT CGC GCA GCT TGC GGA AAC GGG GTA TGG ACC CTG CCG TAC CGT TGT TGC CAC TTG ATG gly arg ala ala cys gly asn gly val trp thr leu pro tyr arg cys cys his leu met 211/71 181/61 TCG TCG CTC TCC ACC CGT CGG GGG GCG AAA GCC ATT CCG ACA CTG GGA TCC TCA AAA CGT ser ser leu ser thr arg arg gly ala lys ala ile pro thr leu gly ser ser lys arg 271/91 CGG CTG AGT GTC TGC AGG GCT CCG GGG AGC AGC CGA TCA TCA CCA TGT ACG AAC TGA ATA arg leu ser val cys arg ala pro gly ser ser arg ser ser pro cys thr asn OPA ile 331/111 301/101 AGT CCC CCC CGC GCG ACT TCC AGA CAT TTG TTG TGG TTT CGG TTG AGG CCG AGG CGA GGC ser pro pro arg ala thr ser arg his leu leu trp phe arg leu arg pro arg arg gly 391/131 TCA TTT CGC AGC AAG CGG TCT CCG GGT CGC AGC ATC GTT GCG GCG ATC GCG GCG CAG TCG ser phe arg ser lys arg ser pro gly arg ser ile val ala ala ile ala ala gln ser 421/141 TCG GAC GAG TCG TCG TCA ACG ACC ACG ATC ser asp glu ser ser ser thr thr thr ile

SEQ ID No.27B

FIGURE 27B
REPLACEMENT SHEET (RULE 26)





	33/11
TAC CAG CAA GAG CCC AGG GCT TCA CAG	GAC CTA AAA GGA GTA GCG CCC ATG GGC TTG ATC
tyr gln gln glu pro arg ala ser gln	asp leu lys gly val ala pro met gly leu ile
63/21	93/31
CAA TTT TCC TTC CGC CCC GTG CAA TAC	CAT CTG CAA GAC CAG CGA CGG CCC GTG GTT GCG
gln phe ser phe arg pro val gln tyr	his leu gln asp gln arg arg pro val val ala
123/41	153/51
GTC GCG CAG CTT GCG GAA ACG GGG TAT	GGA CCC TGC CGT ACC GTT GTT GCC ACT TGA TGT
-	gly pro cys arg thr val val ala thr OPA cys
183/61	
CGT CGC TCT CCA CCC GTC GGG GGG CGA	AAG CCA TTC CGA CAC TGG GAT CCT CAA AAC GTC
	g lys pro phe arg his trp asp pro gln asn val 273/91
243/81	A GCA GCC GAT CAT CAC CAT GTA CGA ACT GAA TAA
GGC TGA GTG TCT GCA GGG CTC CGG GGA	v ala ala asp his his his val arg thr glu OCH
	333/111
303/101	TGT TGT GGT TTC GGT TGA GGC CGA GGC GAG GCT
and has been als are less pro asp ile	e cys cys gly phe gly OPA gly arg gly glu ala
363/121	393/131
CAT TTC GCA GCA AGC GGT CTC CGG GTC	C GCA GCA TCG TTG CGG CGA TCG CGG CGC AGT CGT
his phe ala ala ser gly leu arg val	l ala ala ser leu arg arg ser arg arg ser arg
423/141	
CGG ACG AGT CGT CGT CAA CGA CCA CGA	A TC
arg thr ser arg arg gln arg pro arg	
	-

SEQ ID No.27C

FIGURE 27C

 $\label{thm:mktgtattrrllavlialalpgaavallaepsatgasdpcaasevar tvgsvaksmgdyldshpetnqvmtavlqqqvgpgsvaslkahfeanpk vasdlhalsqpltdlstrcslpisglqaiglmqavqgarr$

SEQ ID No.28

FIGURE 28

GTGGGCAAGC	AGCTAGCCGC	GCTCGCCGCG	CTGGTCGGTG	CGTGCATGCT	CGCAGCCGGA	60
TGCACCAACG	TGGTCGACGG	GACCGCCGTG	GCTGCCGACA	AATCCGGACC	ACTGCATCAG	120
GATCCGATAC	CGGTTTCAGC	GCTTGAAGGG	CTGCTTCTCG	ACTTGAGCCA	GATCAATGCC	180
GCGCTGGGTG	CGACATCGAT	GAAGGTGTGG	TTCAACGCCA	AGGCAATGTG	GGACTGGAGC	240
AAGAGCGTGG	CCGACAAGAA	TTGCCTGGCT	ATCGACGGTC	CAGCACAGGA	AAAGGTCTAT	300
GCCGGCACCG	GGTGGACCGC	TATGCGCGGC	CAACGGCTGG	ATGACAGCAT	CGATGACTCC	360
AAGAAACGCG	ACCACTACGC	CATTCAAGCG	GTCGTCGGCT	TCCCGACCGC	ACATGATGCC	420
GAGGAGTTCT	ACAGCTCCTC	GGTGCAAAGC	TGGAGCAGCT	GCTCGAACCG	CCGGTTTGTC	480
GAAGTCACCC	CCGGACAGGA	CGACGCCGCC	TGGACTGTGG	CTGACGTTGT	CAACGACAAC	540
GGCATGCTCA	GTAGCTCGCA	GGTTCAGGAA	GGCGGCGACG	GATGGACCTG	CCAGCGTGCC	600
	GCAACAACGT					660
TTGGTGGCGA	TTGGCATCGC	TAACCAAATC	GCGGCCAAGG	TTGCTAAGCA	GTAG	714

SEQ ID No.29

FIGURE 29

REPLACEMENT SHEET (RULE 26)





 $\label{thm:mgkqlaalvgacmlaagctnvvdgtavaadksgplhqdpipvftsalegllldlsqinaalgats \\ \texttt{MKVWFNAKAMWDWSKSVADKNCLAIDGPAQEKVYAGTGFTWTAMRGQRLDDSIDDSKKRDHYAIQAVV} \\ \texttt{GFPTAHDAEEFYSSSVQSWSSCSNRFFVEVTFTPGQDDAAWTVADVVNDNGMLSSSQVQEGGDGWTCQ} \\ \texttt{RALTARNNVTIDIVTCAYSQPDLVFTAIGIANQIAAKVAKQ}$

SEQ ID No.30

FIGURE 30

1/1					31/11								
AGG CGA ATA CCC	GCG AGG	GCA GCG	CGA C	CGG	CGG CCC	TGC	CGG	CGC	CGT	GGC	TGC	TGA	ACA
arg arg ile pro a	ala arg	ala ala	arg a	arg	arg pro	cys	arg	arg	arg	gly	cys	OPA	thr
61/21					91/31								
ACA CAT CCC AGC	CGC GCA	CGC TTC	CGG I	TAT	GCG GCA	GGA	TAA	ACG	ACC	CCA	ACA	GCA	CGA
thr his pro ser a	arg ala	arg phe	arg t	tyr	ala ala	gly	OCH	thr	thr	pro	thr	ala	arg
121/41					151/51								
ACA CCA GGA TTG	CGA CAA	CCA AAG	CCC 1	TCG	CGC CTG	GCT	CGA	TTT	CGC	GCG	CAA	CGC	GGC
	CGA CAA arg gln	CCA AAG pro lys	CCC T	TCG ser	CGC CTG arg leu	GCT ala	CGA arg	TTT phe	CGC arg	GCG ala	CAA gln	CGC arg	GGC gly
ACA CCA GGA TTG C thr pro gly leu a 181/61	arg gln	pro lys	pro s	TCG ser	CGC CTG arg leu 211/71	ala	arg	TTT phe	CGC arg	GCG ala	CAA gln	CGC arg	GGC gly
ACA CCA GGA TTG C	arg gln	pro lys	pro s	TCG ser GGG	CGC CTG arg leu 211/71 CGT CGA	ala GAT	arg	TTT phe	CGC arg	GCG ala	CAA gln	CGC arg	GGC gly

SEQ ID No.31A

FIGURE 31A

1/1									31/1	1								
GGC GAA	TAC	CCG	CGA	GGG	CAG	CGC	GAC	GGC	GGC	CCT	GCC	GGC	GCC	GTG	GCT	GCT	GAA	CAA
gly glu	tyr	pro	arq	gly	gln	arg	asp	gly	gly	pro	ala	gly	ala	val	ala	ala	glu	gln
61/21									91/3	31								
CAC ATC	CCA	GCC	GCG	CAC	GCT	TCC	GGT	ATG	CGG	CAG	GAT	AAA	CGA	CCC	CAA	CAG	CAC	GAA
his ile	pro	ala	ala	his	ala	ser	gly	met	arg	gln	asp	lys	arg	pro	gln	gln	his	glu
121/41	-								151,	-								
CAC CAG	GAT	TGC	GAC	AAC	CAA	AGC	CCT	CGC	GCC	TGG	CTC	GAT	TTC	GCG	CGC	AAC	GCG	GCG
his gln	asp	cys	asp	asn	gln	ser	pro	arg	ala	trp	leu	asp	phe	ala	arg	asn	ala	ala
181/61	_	_							211,									
TTC TGC																		
phe cvs	arq	leu	asp	leu	ser	ala	glu	gly	val	glu	ile							

SEQ ID No.31B

FIGURE 31B



31/111/1 GCG AAT ACC CGC GAG GGC AGC GCG ACG GCG GCC CTG CCG GCG CCG TGG CTG CTG AAC AAC ala asn thr arg glu gly ser ala thr ala ala leu pro ala pro trp leu leu asn asn 91/31 61/21 ACA TCC CAG CCG CGC ACG CTT CCG GTA TGC GGC AGG ATA AAC GAC CCC AAC AGC ACG AAC thr ser gln pro arg thr leu pro val cys gly arg ile asn asp pro asn ser thr asn 151/51 121/41 ACC AGG ATT GCG ACA ACC AAA GCC CTC GCG CCT GGC TCG ATT TCG CGC GCA ACG CGG CGT thr arg ile ala thr thr lys ala leu ala pro gly ser ile ser arg ala thr arg arg 211/71 181/61 TCT GCC GCC TCG ATC TCA GCG CGG AGG GCG TCG AGA TC ser ala ala ser ile ser ala arg arg ala ser arg

SEO ID No.31C

FIGURE 31C

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ORF according to Cole et al. (Nature 393:537-544) and containing seq31A
                                        31/11
1/1
taa acg acc cca aca gca cga aca cca gga ttg cga caa cca aag ccc tcg cgc ctg gct
OCH thr thr pro thr ala arg thr pro gly leu arg gln pro lys pro ser arg leu ala
                                        91/31
61/21
cga ttt cgc gcg caa cgc ggc gtt ctg ccg cct cga tct cag cgc gga ggg cgt cga gat
arg phe arg ala gln arg gly val leu pro pro arg ser gln arg gly gly arg arg asp
                                        151/51
121/41
ccc cgg cgt cgt gtt cgt ggc tca tca tct gca tcc tcc ggg ctt ggc cgc gct gac cgg
pro arg arg val arg gly ser ser ser ala ser ser gly leu gly arg ala asp arg
                                        211/71
181/61
cag ccc gac ccc agg cat gcc cag gcc gac ggc gcg ccc cgg ctg ccc ggc ggt gtg cgc
gln pro asp pro arg his ala gln ala asp gly ala pro arg leu pro gly gly val arg
                                        271/91
241/81
gtc gcc ggc gcg ggt gcg gcg gtg ggt cag gac gcc ggc gtc ggc gat gag gtg gtg cgg
val ala gly ala gly ala ala val gly gln asp ala gly val gly asp glu val val arg
                                        331/111
ege ege tte ggt gae ett egt ggt gat gae gte gee ggg aeg eae geg egg etg gee gge
arg arg phe gly asp leu arg gly asp asp val ala gly thr his ala arg leu ala gly
                                        391/131
361/121
ggt gaa gtg cac cag gcg ccc gtc gcg cgc ccg ccc gct cat gcg cgc cgt gac ggt gtc
gly glu val his gln ala pro val ala arg pro pro ala his ala arg arg asp gly val
                                         451/151
421/141
ctt geg eec tte eec ggt gge eac eag eac etc gae gge etg eec gae eag gge geg gtt
leu ala pro phe pro gly gly his gln his leu asp gly leu pro asp gln gly ala val
                                        511/171
481/161
ggc ttc cag cga gat ttg ctc ctg cag cgc gat cag gcg ttc ata gcg ttc ctg cac aac
gly phe gln arg asp leu leu eun arg asp gln ala phe ile ala phe leu his asn
                                         571/191
ggc ttt cgg cag ctg tcc gtc gag ttg cgc ggc cgg tgt ccc ggg ccg ctt gga gta ttg
gly phe arg gln leu ser val glu leu arg gly arg cys pro gly pro leu gly val leu
                                         631/211
601/201
gaa ggt aaa tgc ggc cgc gaa gcg ggc ccg gcg cac cac gtc gag cgt ggc cgc gaa gtc
glu gly lys cys gly arg glu ala gly pro ala his his val glu arg gly arg glu val
                                         691/231
661/221
ctc ttc ggt ctc ccc ggg gaa acc gac gat cag atc ggt ggt aat cgc ggc atg cgg gat
leu phe gly leu pro gly glu thr asp asp gln ile gly gly asn arg gly met arg asp
                                         751/251
721/241
ggc cgc ccg cac gcg ctc gat gat gcc gag gta gcg ctc ggc acg ata gga ccg ccg cat
gly arg pro his ala leu asp asp ala glu val ala leu gly thr ile gly pro pro his
                                         811/271
781/261
cgc gcg cag gat ccg gtc gga tcc gga ctg tag
arg ala gln asp pro val gly ser gly leu AMB
```

SEQ ID No.31F

FIGURE 31F



31/11 1/1 aga ctg gtg tac acg gag acc aag ctg aac tcg gca ttc tcc ttc ggc ggg cct aag tgt arg leu val tyr thr glu thr lys leu asn ser ala phe ser phe gly gly pro lys cys 91/31 cta gtg aag gtc att cag aaa ctg tcg ggc ttg agc atc aac cgg ttc atc gcg att gac leu val lys val ile gln lys leu ser gly leu ser ile asn arg phe ile ala ile asp 151/51 ttc gtc ggt ttc gcg cgg atg gtc gag gcc ctc ggc ggc gtc gag gta tgc agc acc acc phe val gly phe ala arg met val glu ala leu gly gly val glu val cys ser thr thr 211/71 ccg ttg cgg gac tac gaa ctg ggc acg gtg ctg gag cac gcc gga cgc cag gtc att gac pro leu arg asp tyr glu leu gly thr val leu glu his ala gly arg gln val ile asp 271/91 241/81 ggg ccg acc gcg ctg aac tat gtg cgc gct cgc cag gtc acc acc gag agc aat ggc gac gly pro thr ala leu asn tyr val arg ala arg gln val thr thr glu ser asn gly asp 331/111 301/101 tac ggg cgc atc aaa cgc cag cag ttg ttt ttg tcg tcg ctg cgt tcg atg atc tyr gly arg ile lys arg gln gln leu phe leu ser ser leu leu arg ser met ile

SEQ ID No.32A

FIGURE 32A

1/1	31/11	
gac tog tot aca cog aga cca agc tga	act cgg cat tot cot tog gcg ggc cta agt gto	;
asp trp cvs thr arg arg pro ser OPA	thr arg his ser pro ser ala gly leu ser val	
61/21	91/31	
tag tga agg tga ttc aga aac tgt cgg	gct tga gca tca acc ggt tca tcg cga ttg act	-
AMB OPA arg ser phe arg asn cys arg	ala OPA ala ser thr gly ser ser arg leu thr	:
121/41	151/51	
tog tog gtt tog oge gga tgg tog agg	ccc tcg gcg gcg tcg agg tat gca gca cca ccc	2
ser ser val ser arg gly trp ser arg	pro ser ala ala ser arg tyr ala ala pro pro)
181/61	211/71	
cat tac agg act acg aac tag qca cgg	tgc tgg agc acg ccg gac gcc agg tca ttg acg	ş
arg cys gly thr thr asn trp ala arg	cys trp ser thr pro asp ala arg ser leu thr	2
241/81	271/91	
and can acq age tag act atq tag acq	ctc gcc agg tca cca ccg aga gca atg gcg act	-
gly arg pro arg OPA thr met cys ala	leu ala arg ser pro pro arg ala met ala thr	-
301/101	331/111	
acq qqc qca tca aac qcc aqc aqt tgt	ttt tgt cgt cgc tgc tgc gtt cga tga tc	
thr gly ala ser asn ala ser ser cys	phe cys arg arg cys cys val arg OPA	

SEQ ID No.32B

FIGURE 32B



31/11 act ggt gta cac gga gac caa gct gaa ctc ggc att ctc ctt cgg cgg gcc taa gtg tct thr gly val his gly asp gln ala glu leu gly ile leu leu arg arg ala OCH val ser 91/31 agt gaa ggt cat toa gaa act gto ggg ctt gag cat caa cog gtt cat cgc gat tga ctt ser glu gly his ser glu thr val gly leu glu his gln pro val his arg asp OPA leu 151/51 121/41 cgt cgg ttt cgc gcg gat ggt cga ggc cct cgg cgg cgt cga ggt atg cag cac ccc arg arg phe arg ala asp gly arg gly pro arg arg arg gly met gln his his pro 211/71 181/61 gtt gcg gga cta cga act ggg cac ggt gct gga gca cgc cgg acg cca ggt cat tga cgg val ala gly leu arg thr gly his gly ala gly ala arg arg thr pro gly his OPA arg 271/91 241/81 gcc gac cgc gct gaa cta tgt gcg cgc tcg cca ggt cac cac cga gag caa tgg cga cta ala asp arg ala glu leu cys ala arg ser pro gly his his arg glu gln trp arg leu 331/111 301/101 cgg gcg cat caa acg cca gca gtt gtt ttt gtc gtc gct gct gcg ttc gat gat c arg ala his gln thr pro ala val val phe val val ala ala phe asp asp

SEQ ID No.32C

FIGURE 32C

sequence Rv0822c predicted by Cole et al. (Nature 393:537-544) and containing seq 32A

```
31/11
atg agt gac ggc gag agc gcc gcg ccg tgg gca cgg ctc tcc gag tca gca ttc ccc gat
Met ser asp gly glu ser ala ala pro trp ala arg leu ser glu ser ala phe pro asp
                                        91/31
ggt gtt gac cga tgg atc acg gta ccg ccc gcc aca tgg gtg gca gcc cag ggt ccg cgg
gly val asp arg trp ile thr val pro pro ala thr trp val ala ala gln gly pro arg
                                        151/51
gac acc cag aat gtc ggc tgt cat gcc acc ggc gcc gtt agt gtg gcc gat ctg atc gcc
asp thr gln asn val gly cys his ala thr gly ala val ser val ala asp leu ile ala
                                        211/71
agg ctc ggc ccc gct ttt cct gac ctc ccc acg cac cgc cat gtc gcc ccc gaa ccc gag
arg leu gly pro ala phe pro asp leu pro thr his arg his val ala pro glu pro glu
                                        271/91
241/81
cca too ggo ogo ggo cog aag gto cao gao gao goo gao gao cag cag gao aco gag got
pro ser gly arg gly pro lys val his asp asp ala asp asp gln gln asp thr glu ala
                                        331/111
atc gcc atc ccg gcc cac tcg ctc gag ttc ctc tcg gag ctt ccc gac ctc cgg gca gcc
ile ala ile pro ala his ser leu glu phe leu ser glu leu pro asp leu arg ala ala
                                        391/131
aac tat eeg ege gee gae eac gee ege egt gaa eee gag eta eee gge aag eag eta ace
asn tyr pro arg ala asp his ala arg arg glu pro glu leu pro gly lys gln leu thr
                                        451/151
421/141
gga tcg gct cga gtg cgg cca ttg cgg atc cgc cga acg tcg ccc gcg ccc gcc aag cca
gly ser ala arg val arg pro leu arg ile arg arg thr ser pro ala pro ala lys pro
                                        511/171
481/161
gcg ccg aac tcc ggc cgg cgc ccg atg gtg ctg gcc gcg cgc tcg ctg gcg gct ctg ttt
ala pro asn ser gly arg arg pro met val leu ala ala arg ser leu ala ala leu phe
                                        571/191
541/181
gcc gct ctg gcg ttg gcg ctg acc ggc ggg gca tgg cag tgg agc gcg tcg aag aac agc
ala ala leu ala leu ala leu thr gly gly ala trp gln trp ser ala ser lys asn ser
                                         631/211
601/201
cgg ctg aac atg gta agc gcg ctc gac ccg cat tcg ggc gac atc gtc aac ccc agc ggg
arg leu asn met val ser ala leu asp pro his ser gly asp ile val asn pro ser gly
```

SEO ID No.32D

FIGURE 32D

```
691/231
cag cat ggc gac gag aac ttc ttg ctc gtc ggt atg gac tct cgt gcc ggg gcg aac gcc
gln his gly asp glu asn phe leu leu val gly met asp ser arg ala gly ala asn ala
                                      751/251
asn ile gly ala gly asp ala glu asp ala gly gly ala arg ser asp thr val met leu
                                      811/271
781/261
gtc aac att ccg gcc agc cgc gag cgg gtc gtc gcg gtg tcg ttc ccc cgc gac ctg gcg
val asn ile pro ala ser arg glu arg val val ala val ser phe pro arg asp leu ala
                                      871/291
841/281
atc act cca atc caa tgc gag gcg tgg aac ccc gag acc ggt aag tac gga ccc atc tac
ile thr pro ile gln cys glu ala trp asn pro glu thr gly lys tyr gly pro ile tyr
                                      931/311
901/301
gac gag aag acg gga acg atg ggt ccc aga ctg gtg tac acg gag acc aag ctg aac tcg
asp glu lys thr gly thr met gly pro arg leu val tyr thr glu thr lys leu asn ser
                                      991/331
gca ttc tcc ttc ggc ggg cct aag tgt cta gtg aag gtc att cag aaa ctg tcg ggc ttg
ala phe ser phe gly gly pro lys cys leu val lys val ile gln lys leu ser gly leu
                                       1051/351
age ate aac egg tte ate geg att gae tte gte ggt tte geg egg atg gte gag gee ete
ser ile asn arg phe ile ala ile asp phe val gly phe ala arg met val glu ala leu
                                      1111/371
ggc ggc gtc gag gta tgc agc acc ccg ttg cgg gac tac gaa ctg ggc acg gtg ctg
gly gly val glu val cys ser thr thr pro leu arg asp tyr glu leu gly thr val leu
                                       1171/391
1141/381
gag cac gcc gga cgc cag gtc att gac ggg ccg acc gcg ctg aac tat gtg cgc gct cgc
glu his ala gly arg gln val ile asp gly pro thr ala leu asn tyr val arg ala arg
                                       1231/411
1201/401
cag gtc acc acc gag agc aat ggc gac tac ggg cgc atc aaa cgc cag cag ttg ttt ttg
gln val thr thr glu ser asn gly asp tyr gly arg ile lys arg gln gln leu phe leu
                                       1291/431
1261/421
tog tog ctg ctg cgt tog atg atc tog acg gac acc ttg ttc aac ctc agc agg ctc aac
ser ser leu leu arg ser met ile ser thr asp thr leu phe asn leu ser arg leu asn
                                       1351/451
1321/441
aac gtc gtc aac atg ttc atc ggt aac agc tac gtg gac aac gtc aag acc aaa gac ctg
asn val val asn met phe ile gly asn ser tyr val asp asn val lys thr lys asp leu
                                       1411/471
1381/461
gtc gaa ctc ggt cga tcg ttg cag cat atg gcg gcc ggg cac gtc acg ttc gtg acc gtt
val glu leu gly arg ser leu gln his met ala ala gly his val thr phe val thr val
                                       1471/491
1441/481
ccg acc ggt ata acc gac cag aac ggc gac gag ccc ccg cgt acc tcc gac atg aag gcg
pro thr gly ile thr asp gln asn gly asp glu pro pro arg thr ser asp met lys ala
                                       1531/511
 1501/501
ctt ttc acc gcc atc atc gac gac gat ccg ctg ccc ctg gaa aac gat cac aac gcc cag
leu phe thr ala ile ile asp asp pro leu pro leu glu asn asp his asn ala gln
                                       1591/531
arg leu gly asn thr pro ser thr pro pro thr thr lys lys ala pro gln ala gly
                                       1651/551
 1621/541
 ctg acc aac gag att cag cac cag gtt acg acg acc tcg cca aaa gag gtc aca gtg
 leu thr asn glu ile gln his gln gln val thr thr thr ser pro lys glu val thr val
                                       1711/571
 1681/561
 cag gtc tct aac tcg acc ggc cag gcc ggt ttg gcc acc gcc acc gat cag ctc aag
 gln val ser asn ser thr gly gln ala gly leu ala thr thr ala thr asp gln leu lys
                                       1771/591
 1741/581
 cgg aac ggc ttc aac gtg atg gct ccg gac gac tac ccg agt tcg ctg ctg gcc acc aca
 arg asn gly phe asn val met ala pro asp asp tyr pro ser ser leu leu ala thr thr
                                       1831/611
 1801/601
 gtg ttt ttt tcg ccc ggc aac gaa cag gct gcc gcc acc gtg gcc gcc gtg ttc ggc cag
 val phe phe ser pro gly asn glu gln ala ala ala thr val ala ala val phe gly gln
                                       1891/631
 1861/621
 tca aag atc gag cgg gtg acc ggg atc ggc caa ctg gtc cag gtg gtg ctg ggc caa gac
 ser lys ile glu arg val thr gly ile gly gln leu val gln val val leu gly gln asp
```

SEQ ID No.32D (continued 1)

FIGURE 32D (continued 1)



1921/641

ttc agc gcg gtg cgc gct ccc ctg ccg agt ggc tcc acc gtc agc gtg cag ata agc cgc phe ser ala val arg ala pro leu pro ser gly ser thr val ser val gln ile ser arg 1981/661

aac tcc tcc agc cca ccg acc aag ctg ccc gag gac ctg acg gtc acc aac gcc gcc gac asn ser ser ser pro pro thr lys leu pro glu asp leu thr val thr asn ala ala asp 2041/681

acc acc tgc gag tag thr thr cys glu AMB

SEQ ID No.32D (continued 2) FIGURE 32D (continued 2)

ORF according to Cole et al. (Nature 393:537-544) and containing Rv0822c

									31/1	1								
1/1 tag gac				~~~	~~~	200	acc	aca			aca	caa	ctc	tcc	gag	tca	σca	ttc
tag gac AMB asp	atg	agt	gac	ggc	gay	ayc	212	909 ala	nro	trn	ala	ard	leu	ser	alu	ser	ala	phe
	met	ser	asp	GIY	gru	261	ara	aru	91/3	1	424	~~ 9			5			• .
61/21 ccc gat		~++	~~ c	~~~	taa	atc	aca	αta			acc	aca	taa	ata	σca	αcc	caq	aat
pro asp	ggı	gul	gac	ara	trn	ile	thr	val	pro	pro	ala	thr	tro	val	ála	ála	gln	gly
121/41	gry	vai	asp	arg	CLP	110			151/	51			-				-	
cca caa	as c	acc	can	aat	atic	aac	tat	cat	acc	acc	aac	gcc	gtt	agt	gtg	gcc	gat	ctg
pro arg	asn	thr	aln	asn	val	alv	cvs	his	ala	thr	gly	āla	val	ser	val	ala	asp	leu
181/61									211/	71								
ata acc	aσσ	ctc	aac	ccc	gct	ttt	cct	gac	ctc	CCC	acg	cac	cgc	cat	gtc	gcc	CCC	gaa
ile ala	arg	leu	qly	pro	ala	phe	pro	asp	leu	pro	thr	his	arg	his	val	ala	pro	glu
241/81									271/	91								
ccc gag	cca	tcc	ggc	cgc	ggc	ccg	aag	gtc	cac	gac	gac	gcc	gac	gac	cag	cag	gac	acc
pro glu	pro	ser	gly	arg	gly	pro	lys	val	his	asp	asp	ala	asp	asp	gln	gln	asp	thr
301/101									331/	1111								
gag gct	atc	gcc	atc	ccg	gcc	cac	tcg	ctc	gag	ttc	ctc	tcg	gag	ctt	ccc	gac	CEC	cgg
glu ala	ile	ala	ile	pro	ala	his	ser	leu	gLu	phe	Ieu	ser	gru	Leu	pro	asp	Teu	arg
361/121										/131				a+ a		~~~	224	cad
gca gcc	aac	tat	ccg	cgc	gcc	gac	cac	gcc	cgc	cgt	gaa	222	gag	lou	nro	4J v	lvs	aln
ala ala	asn	tyr	pro	arg	ala	asp	nıs	ara	arg	/151	gru	ÞΙΟ	gru	ieu	pro	Q T Y	- 1 2	9111
421/141 cta acc				~~~	~+~	~~~	CC3	++~			cac	cas	acq	t.ca	ccc	aca	ccc	acc
leu thr	gga	tcg	get	cga	gra	ara	nro	1011	ara	ile	ara	arg	thr	ser	pro	ala	pro	ala
	дтХ	ser	ara	ary	vai	ary	pro	ıcu	511	/171	u_9	u_ 9			F		•	
481/161 aag cca	~~~	cca	220	tcc	ממכ	caa	cac	cca			cta	acc	aca	cgc	tcg	ctg	gcg	gct
lys pro	ala	nro	asn	ser	ulv	aro	arg	pro	met	val	leu	ala	ala	arg	ser	leu	ala	ala
541/181									5/1	/ T9T								
cta ttt	acc	act	cta	aca	ttg	gcg	ctg	acc	ggc	ggg	gca	tgg	cag	tgg	agc	gcg	tcg	aag
leu phe	ala	ala	leu	ālā	leu	ālā	leu	thr	gly	gly	ala	trp	gln	trp	ser	ala	ser	lys
601/201									631	/211								
aac agc	cgg	ctg	aac	atg	gta	agc	gcg	ctc	gac	ccg	cat	tcg	gāc	gac	atc	gtc	aac	ccc
asn ser	arg	leu	asn	met	val	ser	ala	leu	asp	pro	his	ser	дтĀ	asp	ше	vaı	asn	pro
661/221										/231			~~~	+-+	cat	acc	aaa	aca
agc ggg	cag	cat	ggc	gac	gag	aac	ttc	ttg	ctc	gtc	ggt	atg	gac	cor	ara	900 ala	999 71 v	909 ala
ser gly		his	gly	asp	gru	asn	pne	reu	. 1eu	vai /251	g _T y	mec	asp	261	ary	ara	9-1	ara
721/241						~~~	. ~~~	~~~			aac	aac	aca	cat	tca	σac	acc	atc
aac gcc asn ala	aat	atc	ggc	gcc	ggc	gac	900	gay	yac	900 ala	al v	ggc dlv	ala	ard	ser	asp	thr	val
		тте	g _T y	аıа	дту	asp	ата	. gru	. asp 811	/271	9-1	9± I		9				
781/261 atg ctg	. ~+-		+			200	cac	· dad	ו כמם	ato	ato	aca	ata	tca	ttc	ccc	cgc	gac
met leu	gcc	adC	ila	nro	gcc	ser	aro	gay alu	ara	val	val	ala	val	ser	phe	pro	arg	asp
met reu	val	ası.		PLO	u La			9	3		/-				-	-	_	

SEQ ID No.32F

FIGURE 32F
REPLACEMENT SHEET (RULE 26)



841/281 871/291	
ctg gcg atc act cca atc caa tgc gag gcg tgg aac ccc gag acc ggt aag tac gga c	cc
leu ala ile thr pro ile gln cys glu ala trp asn pro glu thr gly lys tyr gly p	ro
901/301 931/311 atc tac gac gag aag acg atg ggt ccc aga ctg gtg tac acg gag acc aag c	.+~
ile tyr asp qlu lys thr gly thr met gly pro arg leu val tyr thr glu thr lys l	
961/321 991/331	
aac tog goa tto too tto ggo ggg cot aag tgt cta gtg aag gto att cag aaa ctg t	
asn ser ala phe ser phe gly gly pro lys cys leu val lys val ile gln lys leu s 1021/341 1051/351	er
qqc ttq agc atc aac cgg ttc atc gcg att gac ttc gtc ggt ttc gcg cgg atg gtc g	jag
gly leu ser ile asn arg phe ile ala ile asp phe val gly phe ala arg met val g	,lu
1081/361 1111/371 qcc ctc qgc ggc gtc gag gta tgc agc acc acc ccg ttg cgg gac tac gaa ctg ggc a	
ala leu gly gly val glu val cys ser thr thr pro leu arg asp tyr glu leu gly t	
1141/381 1171/391	
gtg ctg gag cac gcc gga cgc cag gtc att gac ggg ccg acc gcg ctg aac tat gtg c	
val leu glu his ala gly arg gln val ile asp gly pro thr ala leu asn tyr val a 1201/401 1231/411	ırg
gct cgc cag gtc acc acc gag agc aat ggc gac tac ggg cgc atc aaa cgc cag cag t	tg
ala arg gln val thr thr glu ser asn gly asp tyr gly arg ile lys arg gln gln l	.eu
1291/421 1291/431 ttt ttq tcq tcq ctq ctq cgt tcq atq atc tcq acq gac acc ttq ttc aac ctc agc a	aaa
phe leu ser ser leu leu arg ser met ile ser thr asp thr leu phe asn leu ser a	
1321/441 1351/451	
ctc aac aac gtc gtc aac atg ttc atc ggt aac agc tac gtg gac aac gtc aag acc a leu asn asn val val asn met phe ile gly asn ser tyr val asp asn val lys thr l	
1381/461 1411/471	.y.3
gac ctg gtc gaa ctc ggt cga tcg ttg cag cat atg gcg gcc ggg cac gtc acg ttc g	
asp leu val glu leu gly arg ser leu gln his met ala ala gly his val thr phe v	/al
1441/481 1471/491 acc gtt ccg acc ggt ata acc gac cag aac ggc gac gag ccc ccg cgt acc tcc gac a	ita
thr val pro thr gly ile thr asp gln asn gly asp glu pro pro arg thr ser asp m	
1501/501 1531/511	
aag gcg ctt ttc acc gcc atc atc gac gac gat ccg ctg ccc ctg gaa aac gat cac a lys ala leu phe thr ala ile ile asp asp pro leu pro leu glu asn asp his a	
1561/521 1591/531	
gee eag egt etg gge aac aeg eeg teg aee eeg eeg aee aee ae aag aag geg eeg e	
ala gln arg leu gly asn thr pro ser thr pro pro thr thr thr lys lys ala pro g 1621/541 1651/551	ıln
qcq qqt ctq acc aac qaq att caq cac cag cag gtt acg acg acc tcg cca aaa gag g	ıtc
ala gly leu thr asn glu ile gln his gln gln val thr thr thr ser pro lys glu v	7al
1681/561 1711/571 aca gtg cag gtc tct aac tcg acc ggc cag gcc ggt ttg gcc acc acc gcc acc gat c	
thr val gln val ser asn ser thr gly gln ala gly leu ala thr thr ala thr asp g	,ay ;ln
1741/581 1771/591	
etc aag egg aac gge tte aac gtg atg get eeg gae tae eeg agt teg etg etg g	
leu lys arg asn gly phe asn val met ala pro asp asp tyr pro ser ser leu leu a 1801/601 1831/611	ша
acc aca gtg ttt ttt tcg ccc ggc aac gaa cag gct gcc gcc acc gtg gcc gcc gtg t	tc
thr thr val phe phe ser pro gly asn glu gln ala ala ala thr val ala ala val p	he
1861/621 1891/631 ggc cag tca aag atc gag cgg gtg acc ggg atc ggc caa ctg gtc cag gtg gtg ctg g	ממכ:
gly gln ser lys ile glu arg val thr gly ile gly gln leu val gln val val leu g	
1921/641 1951/651	
caa gac ttc agc gcg gtg cgc gct ccc ctg ccg agt ggc tcc acc gtc agc gtg cag a gln asp phe ser ala val arg ala pro leu pro ser gly ser thr val ser val gln i	
1981/661 2011/671	
ago ogo aac too too ago coa cog aco aag otg coo gag gao otg acg gto aco aac g	
ser arg asn ser ser ser pro pro thr lys leu pro glu asp leu thr val thr asn a	ıla
2041/681 gcc gac acc acc tgc gag tag	

SEQ ID 32F (continued 1)

FIGURE 32F (continued 1)

REPLACEMENT SHEET (RULE 26)



1/1 31/11

CGT CAC CTC TGC CAT GGT CCA TCT ACG GTA TCT GCG ACA AGG GCA GCG TCG ATC CCT CGA arg his leu cys his gly pro ser thr val ser ala thr arg ala ala ser ile pro arg 61/21 91/31

CAT GCA GAG TCG GTG TTC GCT TCA CGC GAA CTA GGC GCG CCT AGC CTG GAC GAG TCC CCG his ala glu ser val phe ala ser arg glu leu gly ala pro ser leu asp glu ser pro 121/41 151/51

GGC CGA CAT TCG CCC GAG GCC TTG GCC TCC ATC ACC TAA TTG TGT GCA AAA CCG TAT CTA gly arg his ser pro glu ala leu ala ser ile thr OCH leu cys ala lys pro tyr leu 181/61ATT GAT ACG ATT GCG CAC ATG GCT ATC TGG GAT C ile asp thr ile ala his met ala ile trp asp

SEQ ID No.33A

FIGURE 33A

31/11
GTC ACC TCT GCC ATG GTC CAT CTA CGG TAT CTG CGA CAA GGG CAG CGT CGA TCC CTC GAC val thr ser ala met val his leu arg tyr leu arg gln gly gln arg arg ser leu asp 61/21
ATG CAG AGT CGG TGT TCG CTT CAC GCG AAC TAG GCG CGC CTA GCC TGG ACG AGT CCC CGG met gln ser arg cys ser leu his ala asn AMB ala arg leu ala trp thr ser pro arg 121/41
GCC GAC ATT CGC CCG AGG CCT TGG CCT CCA TCA CCT AAT TGT GTG CAA AAC CGT ATC TAA ala asp ile arg pro arg pro trp pro pro ser pro asn cys val gln asn arg ile OCH 181/61
TTG ATA CGA TTG CGC ACA TGG CTA TCT GGG ATC leu ile arg leu arg thr trp leu ser gly ile

SEQ ID No.33B

FIGURE 33B

1/1

CCG TCA CCT CTG CCA TGG TCC ATC TAC GGT ATC TGC GAC AAG GGC AGC GTC GAT CCC TCG pro ser pro leu pro trp ser ile tyr gly ile cys asp lys gly ser val asp pro ser 61/21

ACA TGC AGA GTC GGT GTT CGC TTC ACG CGA ACT AGG CGC GCC TAG CCT GGA CGA GTC CCC thr cys arg val gly val arg phe thr arg thr arg arg ala AMB pro gly arg val pro 121/41

GGG CCG ACA TTC GCC CGA GGC CTT GGC CTC CAT CAC CTA ATT GTG TGC AAA ACC GTA TCT gly pro thr phe ala arg gly leu gly leu his his leu ile val cys lys thr val ser 181/61

AAT TGA TAC GAT TGC GCA CAT GGC TAT CTG GGA TC asn OPA tyr asp cys ala his gly tyr leu gly

SEQ ID No.33C

FIGURE 33C



sequence Rv1044 predicted by Cole et al. (Nature 393:537-544) and containing seq33A

31/11 1/1 ttg tgt gca aaa ccg tat cta att gat acg att gcg cac atg gct atc tgg gat cgc ctc leu cys ala lys pro tyr leu ile asp thr ile ala his met ala ile trp asp arg leu 91/31 61/21 gte gag gtt gee gee gag caa cat gge tae gte aeg aet ege gat geg ega gae ate gge val glu val ala ala glu gln his gly tyr val thr thr arg asp ala arg asp ile gly 151/51 gtc gac cct gtg cag ctc cgc ctc cta gcg ggg cgc gga cgt ctt gag cgt gtc ggc cga val asp pro val gln leu arg leu leu ala gly arg gly arg leu glu arg val gly arg 211/71 181/61 ggt gtg tac cgg gtg ccc gtg ctg ccg cgt ggt gag cac gac gat ctc gca gcc gca gtg gly val tyr arg val pro val leu pro arg gly glu his asp asp leu ala ala ala val 271/91 241/81 teg tgg act ttg ggg egt ggc gtt atc teg cat gag teg gec ttg geg ett cat gee etc ser trp thr leu gly arg gly val ile ser his glu ser ala leu ala leu his ala leu 331/111 301/101 get gae gtg aac eeg teg ege ate eat ete ace gte eeg ege aac aac eat eeg egt geg ala asp val asn pro ser arg ile his leu thr val pro arg asn asn his pro arg ala 391/131 gee ggg gge gag etg tae ega gtt cae ege ege gae ete eag gea gee eae gte aet teg ala gly gly glu leu tyr arg val his arg arg asp leu gln ala ala his val thr ser 451/151 421/141 gtc gac gga ata ccc gtc acg acg gtt gcg cgc acc atc aaa gac tgc gtg aag acg ggc val asp gly ile pro val thr thr val ala arg thr ile lys asp cys val lys thr gly 511/171 481/161 acg gat cct tat cag ctt cgg gcc gcg atc gag cga gcc gaa gcc gag ggc acg ctt cgt thr asp pro tyr gln leu arg ala ala ile glu arg ala glu ala glu gly thr leu arg 571/191 541/181 cgt ggg tca gca gct gag cta cgc gct gcg ctc gat gag acc act gcc gga tta cgc gct arg gly ser ala ala glu leu arg ala ala leu asp glu thr thr ala gly leu arg ala 601/201 cgg ccg aag cga gca tcg gcg tga arg pro lys arg ala ser ala OPA

SEQ ID No.33D

FIGURE 33D



ORF according to Cole et al. (Nature 393:537-544) and containing Rv1044

```
31/11
1/1
taa ttg tgt gca aaa ccg tat cta att gat acg att gcg cac atg gct atc tgg gat cgc
OCH leu cys ala lys pro tyr leu ile asp thr ile ala his met ala ile trp asp arg
                                        91/31
ctc gtc gag gtt gcc gcc gag caa cat ggc tac gtc acg act cgc gat gcg cga gac atc
leu val glu val ala ala glu gln his gly tyr val thr thr arg asp ala arg asp ile
                                        151/51
121/41
ggc gtc gac cct gtg cag ctc cgc ctc cta gcg ggg cgc gga cgt ctt gag cgt gtc ggc
gly val asp pro val gln leu arg leu leu ala gly arg gly arg leu glu arg val gly
                                        211/71
cga ggt gtg tac cgg gtg ccc gtg ctg ccg cgt ggt gag cac gac gat ctc gca gcc gca
arg gly val tyr arg val pro val leu pro arg gly glu his asp asp leu ala ala
                                        271/91
241/81
gtg tcg tgg act ttg ggg cgt ggc gtt atc tcg cat gag tcg gcc ttg gcg ctt cat gcc
val ser trp thr leu gly arg gly val ile ser his glu ser ala leu ala leu his ala
                                        331/111
ctc gct gac gtg aac ccg tcg cgc atc cat ctc acc gtc ccg cgc aac aac cat ccg cgt
leu ala asp val asn pro ser arg ile his leu thr val pro arg asn asn his pro arg
                                         391/131
361/121
gcg gcc ggg ggc gag ctg tac cga gtt cac cgc cgc gac ctc cag gca gcc cac gtc act
ala ala gly gly glu leu tyr arg val his arg arg asp leu gln ala ala his val thr
                                         451/151
421/141
teg gte gae gga ata eee gte acg acg gtt geg ege ace ate aaa gae tge gtg aag acg
ser val asp gly ile pro val thr thr val ala arg thr ile lys asp cys val lys thr
                                         511/171
481/161
ggc acg gat cct tat cag ctt cgg gcc gcg atc gag cga gcc gaa gcc gag ggc acg ctt
gly thr asp pro tyr gln leu arg ala ala ile glu arg ala glu ala glu gly thr leu
                                         571/191
541/181
cgt cgt ggg tca gca gct gag cta cgc gct gcg ctc gat gag acc act gcc gga tta cgc
arg arg gly ser ala ala glu leu arg ala ala leu asp glu thr thr ala gly leu arg
601/201
gct cgg ccg aag cga gca tcg gcg tga
ala arg pro lys arg ala ser ala OPA
```

SEQ ID No.33F

FIGURE 33F

```
31/11
1/1
ATC CAA CCT GCT GGG CCT GCG CCT TCG AAT CGA CGG CCA GGC CAC CGC TCG CTG CCG GCA
ile gln pro ala gly pro ala pro ser asn arg pro gly his arg ser leu pro ala
                                        91/31
61/21
ACA ACA CCT GGA ATG GGG ACC TTT TCG GTG TTG CTG GTA ACC GGG ACA ACC GGC ACC
thr thr pro gly met gly thr phe ser val leu leu val thr gly thr thr gly thr thr
                                        151/51
121/41
CCT CGG TCG AGA CGT ATC GCG GCA GCG TTG GCC CTG TCG TTG CTG ACA ATT ACC GCT GGC
pro arg ser arg arg ile ala ala ala leu ala leu ser leu leu thr ile thr ala gly
                                        211/71
181/61
CGC CGC ATA TTT GCC GCG CTG CCG CGG GCC GGA TC
arg arg ile phe ala ala leu pro arg ala gly
```

SEQ ID No.34A

FIGURE 34A



1/1
TCC AAC CTG CTG GGC CTG CGC CTT CGA ATC GAC GGC CAG GCC ACC GCT CGC TGC CAA
ser asn leu leu gly leu arg leu arg ile asp gly gln ala thr ala arg cys arg gln
61/21
CAA CAC CTG GAA TGG GGA CCT TTT CGG TGT TGC TGG TAA CCG GGA CAA CCG GCA CCA CGC
gln his leu glu trp gly pro phe arg cys cys trp OCH pro gly gln pro ala pro arg
121/41
CTC GGT CGA GAC GTA TCG CGG CAG CGT TGG CCC TGT CGT TGC TGA CAA TTA CCG CTG GCC
leu gly arg asp val ser arg gln arg trp pro cys arg cys OPA gln leu pro leu ala
181/61
GCC GCA TAT TTG CCG CGC TGC CGC GGG CCG GAT C
ala ala tyr leu pro arg cys arg gly pro asp

SEQ ID No.34B

FIGURE 34B

1/1
GAT CCA ACC TGC TGG GCC TGC GCC TTC GAA TCG ACG GCC AGG CCA CCG CTC GCT GCC GGC asp pro thr cys trp ala cys ala phe glu ser thr ala arg pro pro leu ala ala gly 91/31
AAC AAC ACC TGG AAT GGG GAC CTT TTC GGT GTT GCT GGT AAC CGG GAC AAC CGG CAC CAC asn asn thr trp asn gly asp leu phe gly val ala gly asn arg asp asn arg his his 121/41
GCC TCG GTC GAG ACG TAT CGC GGC AGC GTT GCC GTC GTC GTT GCT GAC AAT TAC CGC TGG ala ser val glu thr tyr arg gly ser val gly pro val val ala asp asn tyr arg trp 181/61
CCG CCG CAT ATT TGC CGC GCC GCC GCC GGC CGG ATC pro pro his ile cys arg ala ala ala gly arg ile

SEQ ID No.34C

FIGURE 34C



ORF according to Cole et al. (Nature 393:537-544) containing seq34A

31/11 tag deg dag gge det geg get agg ege gge egg tge egt tgg deg egg egg caa teg atg AMB pro gln gly pro ala ala arg arg gly arg cys arg trp pro arg arg gln ser met 61/21 91/31 ttg cag cag tta caa cgc caa atg gag tct gag cgc atc gtc gag ttc gat cag ctc ggc leu gln gln leu gln arg gln met glu ser glu arg ile val glu phe asp gln leu gly 151/51 agg gga gac gtt gcg cag cga cgg atc caa cct gct ggg cct gcg cct tcg aat cga cgg arg gly asp val ala gln arg arg ile gln pro ala gly pro ala pro ser asn arg arg 181/61 211/71 eca gge cae ege teg etg ecg gea aca aca ect gga atg ggg ace ttt teg gtg ttg etg pro gly his arg ser leu pro ala thr thr pro gly met gly thr phe ser val leu leu 271/91 241/81 gta acc ggg aca acc ggc acc acg cct cgg teg aga cgt atc gcg gca gcg ttg gcc ctg val thr gly thr thr gly thr thr pro arg ser arg arg ile ala ala leu ala leu 301/101 331/111 teg ttg etg aca att ace get gge ege ege ata ttt gee geg etg eeg egg gee gga tee ser leu leu thr ile thr ala gly arg arg ile phe ala ala leu pro arg ala gly ser 391/131 agg teg acc tge cag ate tea eeg ege age ate tae gee gtt ege tge aaa eeg eeg act arg ser thr cys gln ile ser pro arg ser ile tyr ala val arg cys lys pro pro thr 451/151 421/141 geg aeg gea gge eea ete tet tgg eat geg tee aat get geg aeg tee teg gta gae aag ala thr ala gly pro leu ser trp his ala ser asn ala ala thr ser ser val asp lys 511/171 481/161 ctc acg ctt ggc ttc atg ccg cag tcc tac cca tgt agt aac aga tag leu thr leu gly phe met pro gln ser tyr pro cys ser asn arg AMB

SEQ ID No.34F

FIGURE 34F

1/1 31/11 CAG TCT GTC GGC AAG GAG GGA CGC ATG CCA CTC TCC GAT CAT GAG CAG CGG ATG CTT GAC gln ser val gly lys glu gly arg met pro leu ser asp his glu gln arg met leu asp 91/31 CAG ATC GAG AGC GCT CTC TAC GCC GAA GAT CCC AAG TTC GCA TCG AGT GTC CGT GGC GGG gln ile glu ser ala leu tyr ala glu asp pro lys phe ala ser ser val arg gly gly 121/41 151/51 GGC TTC CGC GCA CCG ACC GCG CGG CGC CTG CAG GGC GCG GCG TTG TTC ATC ATC GGT gly phe arg ala pro thr ala arg arg leu gln gly ala ala leu phe ile ile gly 211/71 CTG GGG ATG TTG GTT TCC GGC GTG GCG TTC AAA GAG ACC ATG ATC GGA AGT TTC CCG ATA leu gly met leu val ser gly val ala phe lys glu thr met ile gly ser phe pro ile 241/81 271/91 CTC AGC GTT TTC GGT TTT GTC GTG ATG TTC GGT GGT GTG GTG TAT GCC ATC ACC GGT CCT leu ser val phe gly phe val val met phe gly gly val val tyr ala ile thr gly pro 331/111 301/101 CGG TTG TCC GGC AGG ATG GAT CGT GGC GGA TCG GCT GCT GGG GCT TCG CGC CAG CGT CGT arg leu ser gly arg met asp arg gly gly ser ala ala gly ala ser arg gln arg arg 361/121 391/131 ACC AAG GGG GCC GGG GGC TCA TTC ACC AGC CGT ATG GAA GAT C thr lys gly ala gly gly ser phe thr ser arg met glu asp

SEQ ID No.35A

FIGURE 35A
REPLACEMENT SHEET (RULE 26)



31/11 1/1 GAC AGT CTG TCG GCA AGG AGG GAC GCA TGC CAC TCT CCG ATC ATG AGC AGC GGA TGC TTG asp ser leu ser ala arg arg asp ala cys his ser pro ile met ser ser gly cys leu 91/31 61/21 ACC AGA TCG AGA GCG CTC TCT ACG CCG AAG ATC CCA AGT TCG CAT CGA GTG TCC GTG GCG thr arg ser arg ala leu ser thr pro lys ile pro ser ser his arg val ser val ala 151/51 121/41 GGG GCT TCC GCG CAC CGA CCG CGC GGC GCC TGC AGG GCG CGG CGT TGT TCA TCA gly ala ser ala his arg pro arg gly gly ala cys arg ala arg arg cys ser ser ser 211/71 181/61 GTC TGG GGA TGT TGG TTT CCG GCG TGG CGT TCA AAG AGA CCA TGA TCG GAA GTT TCC CGA val trp gly cys trp phe pro ala trp arg ser lys arg pro OPA ser glu val ser arg 271/91 241/81 TAC TCA GCG TTT TCG GTT TTG TCG TGA TGT TCG GTG GTG TGG TGT ATG CCA TCA CCG GTC tyr ser ala phe ser val leu ser OPA cys ser val val trp cys met pro ser pro val 331/111 301/101 CTC GGT TGT CCG GCA GGA TGG ATC GTG GCG GAT CGG CTG CTG GGG CTT CGC GCC AGC GTC leu gly cys pro ala gly trp ile val ala asp arg leu leu gly leu arg ala ser val 391/131 361/121 GTA CCA AGG GGG CCG GGG GCT CAT TCA CCA GCC GTA TGG AAG ATC val pro arg gly pro gly ala his ser pro ala val trp lys ile

SEQ ID No.35B

FIGURE 35B

31/11 1/1 ACA GTC TGT CGG CAA GGA GGG ACG CAT GCC ACT CTC CGA TCA TGA GCA GCG GAT GCT TGA thr val cys arg gln gly gly thr his ala thr leu arg ser OPA ala ala asp ala OPA 91/31 CCA GAT CGA GAG CGC TCT CTA CGC CGA AGA TCC CAA GTT CGC ATC GAG TGT CCG TGG CGG pro asp arg glu arg ser leu arg arg ser gln val arg ile glu cys pro trp arg 151/51 121/41 GGG CTT CCG CGC ACC GAC CGC GCG GCG GCG CCT GCA GGG CGC GGC GTT GTT CAT CAT gly leu pro arg thr asp arg ala ala pro ala gly arg gly val val his his arg 211/71 181/61 TCT GGG GAT GTT GGT TTC CGG CGT GGC GTT CAA AGA GAC CAT GAT CGG AAG TTT CCC GAT ser gly asp val gly phe arg arg gly val gln arg asp his asp arg lys phe pro asp 271/91 241/81 ACT CAG CGT TTT CGG TGT CGT GAT GTT CGG TGG TGT GGT GTA TGC CAT CAC CGG TCC thr gln arg phe arg phe cys arg asp val arg trp cys gly val cys his his arg ser 331/111 301/101 TCG GTT GTC CGG CAG GAT GGA TCG TGG CGG ATC GGC TGC TGG GGC TTC GCG CCA GCG TCG ser val val arg gln asp gly ser trp arg ile gly cys trp gly phe ala pro ala ser 391/131 361/121 TAC CAA GGG GGC CGG GGG CTC ATT CAC CAG CCG TAT GGA AGA TC tyr gln gly gly arg gly leu ile his gln pro tyr gly arg

SEQ ID No.35C

FIGURE 35C



sequence Rv2169c predicted by Cole et al. (Nature 393:537~544) and partially containing seq35A

31/11 atg cca ctc tcc gat cat gag cag cgg atg ctt gac cag atc gag agc gct ctc tac gcc Met pro leu ser asp his glu gln arg met leu asp gln ile glu ser ala leu tyr ala 61/21 91/31 gaa gat eee aag tte gea teg agt gte egt gge ggg gge tte ege gea eeg aee geg egg qlu asp pro lys phe ala ser ser val arg gly gly gly phe arg ala pro thr ala arg 151/51 121/41 egg ege etg eag gge geg ttg tte ate ate ggt etg ggg atg ttg gtt tee gge gtq arg arg leu gln gly ala ala leu phe ile ile gly leu gly met leu val ser gly val 211/71 181/61 gcg ttc aaa gag acc atg atc gga agt ttc ccg ata ctc agc gtt ttc ggt ttt gtc gtg ala phe lys glu thr met ile gly ser phe pro ile leu ser val phe gly phe val val 271/91 241/81 atg ttc ggt ggt gtg gtg tat gcc atc acc ggt cct cgg ttg tcc ggc agg atg gat cgt met phe gly gly val val tyr ala ile thr gly pro arg leu ser gly arg met asp arg 331/111 301/101 gge gga teg get get ggg get teg ege eag egt egt ace aag ggg gee ggg gge tea tte gly gly ser ala ala gly ala ser arg gln arg arg thr lys gly ala gly gly ser phe 391/131 361/121 acc agc cgt atg gaa gat cgg ttc cgg cgc cgc ttc gac gag taa thr ser arg met glu asp arg phe arg arg phe asp glu OCH

SEQ ID No.35D

FIGURE 35D

ORF according to Cole et al. (Nature 393:537-544) and containing Rv2169c

31/11 1/1tga cag tot gtc ggc aag gag gga cgc atg cca ctc tcc gat cat gag cag cgg atg ctt OPA gln ser val gly lys glu gly arg met pro leu ser asp his glu gln arg met leu 61/21 91/31 gac cag atc gag agc gct ctc tac gcc gaa gat ccc aag ttc gca tcg agt gtc cgt ggc asp gln ile glu ser ala leu tyr ala glu asp pro lys phe ala ser ser val arg gly 121/41 151/51 ggg ggc ttc cgc gca ccg acc gcg cgg cgc ctg cag ggc gcg gcg ttg ttc atc atc gly gly phe arg ala pro thr ala arg arg leu gln gly ala ala leu phe ile ile 181/61 211/71 ggt ctg ggg atg ttg gtt tcc ggc gtg gcg ttc aaa gag acc atg atc gga agt ttc ccg gly leu gly met leu val ser gly val ala phe lys glu thr met ile gly ser phe pro 241/81 271/91 ata etc age gtt ttc ggt ttt gtc gtg atg ttc ggt gtg gtg tat gcc atc acc ggt ile leu ser val phe gly phe val val met phe gly gly val val tyr ala ile thr gly 301/101 331/111 cet egg ttg tee gge agg atg gat egt gge gga teg get ggt ggt teg ege eag egt pro arg leu ser gly arg met asp arg gly gly ser ala ala gly ala ser arg gln arg 391/131 361/121 cgt acc aag ggg gcc ggg ggc tca ttc acc agc cgt atg gaa gat cgg ttc cgg cgc cgc arg thr lys gly ala gly gly ser phe thr ser arg met glu asp arg phe arg arg 421/141 ttc gac gag taa phe asp glu OCH

SEO ID 35F

FIGURE 35F



1/1 31/11 GAC CTG GGA CGA AGA CGA CGG CAG CCG CAA TCA GAT CTA CCC GGT CCT GGT CAA CGT asp leu gly arg arg arg gln gln pro gln ser asp leu pro gly pro gly gln arg 91/31 CAA TGG ACA CCC GAC TAC GGT GCG CCT GCG CGC CTC GAC AAT GCG CGG TTC CTG TTG CCC gln trp thr pro asp tyr gly ala pro ala arg leu asp asn ala arg phe leu leu pro 151/51 121/41 GTG GTC GGA GTG CCA CCC GAC CAG GCC ACC GAC TTC GGC TCC GCT GTT GCA CCA GAA ACG val val gly val pro pro asp gln ala thr asp phe gly ser ala val ala pro glu thr 211/71 ACG GCG CCG GTC TGG ATC ACC ATG CTG TGG CCG CTG GCC GAC CGG CCC CGG TTG GCC CCC thr ala pro val trp ile thr met leu trp pro leu ala asp arg pro arg leu ala pro 241/81 271/91 GGG GCA CCC GGT GGC ACC GTT CCC GTC CGG CTG GTC GAC GAC GAC CTG GCA AAC TCG CTG gly ala pro gly gly thr val pro val arg leu val asp asp leu ala asn ser leu 331/111 GCC AAC GGC GGC CTG GAC ATC CTC CTG TCG GCG GCC GAG TTC GCC ACC AAC CGG GAA ala asn gly gly arg leu asp ile leu leu ser ala ala glu phe ala thr asn arg glu 391/131 GTC GAC CCC GAC GGC GCC GTC GGC CGA GCG CTG TGC CTG GCC ATC GAC CCA GAT C val asp pro asp gly ala val gly arg ala leu cys leu ala ile asp pro asp

SEQ ID No.36A

FIGURE 36A

31/11 1/1 ACC TGG GAC GAC GAC GGC AGC AGC CGC AAT CAG ATC TAC CCG GTC CTG GTC AAC GTC thr trp asp glu asp asp gly ser ser arg asn gln ile tyr pro val leu val asn val 91/31 61/21 AAT GGA CAC CCG ACT ACG GTG CGC CTG CGC GGC TCG ACA ATG CGC GGT TCC TGT TGC CCG asn gly his pro thr thr val arg leu arg gly ser thr met arg gly ser cys cys pro 151/51 121/41 TGG TCG GAG TGC CAC CCG ACC AGG CCA CCG ACT TCG GCT CCG CTG TTG CAC CAG AAA CGA trp ser glu cys his pro thr arg pro pro thr ser ala pro leu leu his gln lys arg 211/71 181/61 CGG CGC CGG TCT GGA TCA CCA TGC TGT GGC CGC TGG CCG ACC GGC CCC GGT TGG CCC CCG arg arg ser gly ser pro cys cys gly arg trp pro thr gly pro gly trp pro pro 241/81 271/91 GGG CAC CCG GTG GCA CCG TTC CCG TCC GGC TGG TCG ACG ACG ACC TGG CAA ACT CGC TGG gly his pro val ala pro phe pro ser gly trp ser thr thr trp gln thr arg trp 301/101 331/111 CCA ACG GCG GCC GGC TGG ACA TCC TCC TGT CGG CGG CCG AGT TCG CCA CCA ACC GGG AAG pro thr ala ala gly trp thr ser ser cys arg arg pro ser ser pro pro thr gly lys 391/131 361/121 TCG ACC CCG ACG GCG CCG TCG GCC GAG CGC TGT GCC TGG CCA TCG ACC CAG ATC ser thr pro thr ala pro ser ala glu arg cys ala trp pro ser thr gln ile

SEQ ID No.36B

FIGURE 36B



1/1 31/11 CCT GGG ACG AAG ACG GCA GCA GCC GCA ATC AGA TCT ACC CGG TCC TGG TCA ACG TCA pro gly thr lys thr thr ala ala ala ile arg ser thr arg ser trp ser thr ser 91/31 61/21 ATG GAC ACC CGA CTA CGG TGC GCC TGC GCG GCT CGA CAA TGC GCG GTT CCT GTT GCC CGT met asp thr arg leu arg cys ala cys ala ala arg gln cys ala val pro val ala arg 121/41 151/51 GGT CGG AGT GCC ACC CGA CCA GGC CAC CGA CTT CGG CTC CGC TGT TGC ACC AGA AAC GAC gly arg ser ala thr arg pro gly his arg leu arg leu arg cys cys thr arg asn asp 211/71 GGC GCC GGT CTG GAT CAC CAT GCT GTG GCC GCT GGC CGA CCG GCC CCG GTT GGC CCC CGG gly ala gly leu asp his his ala val ala ala gly arg pro ala pro val gly pro arg 241/81 271/91 GGC ACC CGG TGG CAC CGT TCC CGT CCG GCT GGT CGA CGA CGA CCT GGC AAA CTC GCT GGC gly thr arg trp his arg ser arg pro ala gly arg arg pro gly lys leu ala gly 331/111 301/101 CAA CGG CGG CCG GCT GGA CAT CCT CCT GTC GGC GGC CGA GTT CGC CAC CAA CCG GGA AGT gln arg arg pro ala gly his pro pro val gly gly arg val arg his gln pro gly ser 361/121 391/131 CGA CCC CGA CGG CGC CGT CGG CCG AGC GCT GTG CCT GGC CAT CGA CCC AGA TC arg pro arg arg arg arg pro ser ala val pro gly his arg pro arg

SEQ ID No.36 C

FIGURE 36C

Coding sequence Rv3909 predicted by Cole et al., 1998 (Nature 393 537-544) containing Seq 36A

1/1 31/11 GTG ACC GCA CTG CAA CTC GGC TGG GCC GCT TTG GCG CGC GTC ACC TCA GCG ATC GGC GTC met thr ala leu gln leu gly trp ala ala leu ala arg val thr ser ala ile gly val 61/21 91/31 GTG GCC GGC CTC GGG ATG GCG CTC ACG GTA CCG TCG GCG GCA CCG CAC GCG CTC GCA GGC val ala gly leu gly met ala leu thr val pro ser ala ala pro his ala leu ala gly 151/51 GAG CCC AGC CCG ACG CCT TTT GTC CAG GTC CGC ATC GAT CAG GTG ACC CCG GAC GTG GTG glu pro ser pro thr pro phe val gln val arg ile asp gln val thr pro asp val val 181/61 211/71 ACC ACT TCC AGC GAA CCC CAT GTC ACC GTC AGC GGA ACG GTG ACC AAT ACC GGT GAC CGC thr thr ser ser glu pro his val thr val ser gly thr val thr asn thr gly asp arg 271/91 CCA GTC CGC GAT GTG ATG GTC CGG CTT GAG CAC GCC GCC GCG GTC ACG TCG TCA ACG GCG pro val arg asp val met val arg leu glu his ala ala ala val thr ser ser thr ala 301/101 331/111 TTA CGC ACC TCG CTC GAC GGC GGC ACC GAC CAG TAC CAG CCG GCC GCG GAC TTC CTC ACG leu arg thr ser leu asp gly gly thr asp gln tyr gln pro ala ala asp phe leu thr

SEQ ID No.36D

FIGURE 36D



361/121	222	~ 7	cm »	~~~	000	000	~ 7 7	C T C	391/			B G G	ama	mac	caa	000	~~~	~~~
GTC GCC																		
val ala	pro	gru	reu	asp	arg	дтА	gin	gru	451/		pne	tnr	Ten	ser	aıa	pro	leu	arg
421/141 TCG CTG	7,00	מככ	ccc	ጥሮር	መመረ	CCC	CTTC	አአሮ	-		GGG	איייר	ሞልሮ	ccc	CTC	CTC	CTC	7.7.0
ser leu																		
481/161	CIIL	ary	pro	261	ıcu	ата	Val	asii	511/	_	9±1	110	CYL	pro	val	ıcu	var	asii
GTC AAT	GGG	ACA	ccc	GAC	TAC	GGT	GCG	ССТ			СТС	GAC	ААТ	GCG	CGG	ТТС	CTG	TTG
val asn																		
541/181	9-1	J	P-U	~-F	-1-	5-1		r	571/						9	<i></i>		
CCC GTG	GTC	GGA	GTG	CCA	CCC	GAC	CAG	GCC	ACC	GAC	TTC	GGC	TCC	GCT	GTT	GCA	CCA	GAA
pro val	val	gly	val	pro	pro	asp	gln	ala	thr	asp	phe	gly	ser	ala	val	ala	pro	glu
601/201									631/	211								
ACG ACG																		
thr thr	ala	pro	val	trp	ile	thr	met	leu	_		leu	ala	asp	arg	pro	arg	leu	ala
661/221									691/									
CCC GGG																		
pro gly	ala	pro	gly	gly	thr	vai	pro	val			val	asp	asp	asp	leu	ala	asn	ser
721/241 CTG GCC	330	CCC	ccc	000	cmc	CAC	אוווע	cmc	751/		ccc	ccc	CAC	mmc	ccc	700	77.7	CCC
leu ala																		
781/261	asii	дтĀ	gry	arg	reu	asp	116	reu	811/		ата	ала	gru	pne	ата	CIII		ary
GAA GTC	GAC	CCC	GAC	GGC	GCC	GTC	GGC	CGA			TGC	CTG	GCC	АТС	GAC	CCA	GAT	СТА
glu val																		
841/281	abp	P		9-1			51		871/		2					-	-	
CTC ATC	ACC	GTC	AAT	GCG	ATG	ACC	GGC	GGC	TAC	GTC	GTG	TCC	GAC	TCG	CCC	GAC	GGG	GCC
leu ile	thr	val	asn	ala	met	thr	gly	gly	tyr	val	vaŀ	ser	asp	ser	pro	asp	gly	ala
901/301									931/	311								
GCT CAA																		
ala gln	leu	pro	gly	thr	pro	thr	his	pro			gly	gln	ala	ala	ala	ser	ser	trp
961/321									991/			- ~~	~~~	~~~	~~~		~~~	~~~
CTG GAT																		
leu asp 1021/34	_	leu	arg	tnr	reu	vaı	nıs	arg	1051			CILL	pro	reu	bro	pne	ala	gru
GCC GAC		CΔΨ	CCT	ጥጥር	CAG	CGG	Curu	тαа				CTG	AGC	GCG	ATC	GCA	ACC	ልጥሮ
ala asp																		
1081/36		чор	4.24	104	9	9			1111									
AGC CCC		GAC	ATC	GTC	GAC	CGC	ATC	CTG	GAT	GTC	AGC	TCC	ACC	CGC	GGC	GCA	ACC	GTG
ser pro	ala	asp	ile	val	asp	arg	ile	leu	asp	val	ser	ser	thr	arg	gly	ala	thr	val
1141/38		_							1171	-								
CTG CCC																		
leu pro		gly	pro	leu	thr	gly	arg	ala				leu	ser	thr	his	gly	asn	thr
1201/40									1231				~~~			~~~		~~~
GTT GCC																		
val ala		ala	ala	ala	asp	phe	ser	pro	-	_	-	дти	дтĀ	ser	ser	g⊥n	11e	дтА
1261/42: TCC GCG		mm v	ccc	CCT	N.C.C	ccc	ccc	CCC	1291			ccc	ccc	CTC	CT A	GCG	ccc	CCG
ser ala																		
1321/44		ieu	Pro	ата	CILL	ara	PLO	arg	1351			Pro	arg	Val	Val	ulu	ara	Pro
TTT GAT		GCG	GTC	GGG	GCC	GCG	CTG	GCC				ACA	AAC	CCG	ACC	GTT	CCT	ACC
phe asp																		
1381/46				J = 1					1411					-			-	
TAT CTA	GAT	CCC	TCG	TTG	TTC	GTT	CGG	ATC	GCG	CAT	GAA	TCG	ATC	ACC	GCG	CGC	CGC	CAG
tyr leu	asp	pro	ser	leu	phe	val	arg	ile	ala	his	glu	ser	ile	thr	ala	arg	arg	gln

SEQ ID No.36D (continued 1)

FIGURE 36D (continued 1)

REPLACEMENT SHEET (RULE 26)



3 4 4 3 / 4 0 3					1471/4	491								
1441/481 GAC GCC TTG GGC	CCA ATC	כדה דהה	CGC				CG 2	AAT	GCC (GCG	CCC	CGT .	ACC	CAA
asp ala leu gly	ala met	leu tro	arg	ser	leu q	lu p	ro	asn	ala a	ala	pro	arg	thr	gln
1501/501					1531/	511								
ARC CRC CRC CCC	CCG GCG	TCG TGG	AGC	CTG	GCC A	GC G	AC	GAC	GCG '	CAG	GTC	ATC	CTG	ACC
ile leu val pro	pro ala	ser trp	ser	leu	ala s	er a	sp .	asp	ala	gln	val	ile	leu	thr
1561/501					1591/	531								
CCC CTC CCC ACC	GCC ATC	CGG TCT	GGC	CTG	GCC G	TG C	CCG	CGA	CCA	CTA	CCG	GCG	GTG	ATC
ala leu ala thr	ala ile	arg ser	gly	leu	ala v	al p	ro	arg	pro	leu	pro	ala	val	ile
1621/541					1651/	551								
GCT GAC GCC GCG	GCC CGC	ACC GAG	CCA	CCG	GAA C	.cc c	CCG	GGC	GCT	TAC	AGC	GCC	GCT	CGC
ala asp ala ala	ala arg	thr glu	pro	pro	glu p	ro p	oro	gly	ala	tyr	ser	ala	ala	arg
1681/561					1711/			~- ~	0mm	000	222	OTT 3	mcc	770
GGC CGG TTC AAT	GAC GAC	ATC ACC	ACG	CAG	ATC G	GC 0	∃GG ¬	CAG	GTT	GCC	066	CTA	166	AAG
gly arg phe asn	asp asp	ile thr	thr	gln	ile g	TÀ d	ЯŢХ	gın	vaı	а⊥а	arg	reu	crb	туѕ
1741/581				~~~	1771/	291	7000	700	ccc	CmC	CNC	ጥ አ ር	אככ	CCA
CTG ACC TCG GCG	TTG ACC	ATC GAT	GAC	CGC	ACC G	.1., 1	J Cu	thr	410	GIG Val	aln	tur	thr	ala
leu thr ser ala	leu thr	ile asp	asp	arg	1831/	'611	Leu	CIII	9×1	Val	9	~ <u>, </u>	0	424
1801/601 CCA CTA CGC GAG	C3 C 3 MC	ששכ ככנ	cce	ርጥር			rcg	СТА	CCA	ccc	GAT	ACC	CGC	AAC
pro leu arg glu	GAC AIG	leu arc	ele:	leu	ser a	ıln :	ser	leu	pro	pro	asp	thr	arq	asn
	asp mec	Ted are	, ara	104	1891/	631			.	-	-			
1861/621 GGG CTG GCC CAG	CAG CGG	CTG GCC	GTC	GTT	GGA A	AAG A	ACG	ATC	GAC	GAT	CTT	TTC	GGC	GCG
gly leu ala gln	aln ara	leu ala	val	val	alv 1	vs '	thr	ile	asp	asp	leu	phe	gly	ala
1021/6/1					1951/	651								
CEC ACC AEC CEC	AAC CCG	GGC GGG	TCC	TAC	ACT C	CTG (GCC	ACC	GAG	CAC	AGT	CCG	CTG	CCG
val thr ile val	asn pro	gly gly	ser	tyr	thr 1	Leu .	ala	thr	glu	his	ser	pro	leu	pro
1091/661					2011/	(0/T								
TTC CCC CTC CAT	AAT GGC	CTC GC	GTG	CCA	ATC C	CGG	GTC	CGG	CTA	CAG	GTC	GAT	GCT	CCG
leu ala leu his	asn gly	leu ala	a val	pro	ile a	arg	val	arg	leu	gln	val	asp	a⊥a	pro
2041/681					2071/	/691		~~~		000	m 7 C	CITIC.	ccc	Cmx
CCC GGG ATG ACG	GTG GCC	GAT GT	GGT	CAG	ATC G	GAG	CTA	CCG		~1	TAC	1011	nra	LON
pro gly met thr	val ala	asp val	r ara	gin	11e 9	gru 7711	reu	pro	pro	gry	CAT	ieu	PIC	ieu
2101/701				~~~	2131/	/ /ll	~~~	СШС	CAC	CTC	ሞሮር	ርጥር	CGG	ACC
CGA GTA CCA ATC	GAG GTG	AAC TT	CACA	CAG	066	JTT m	212	7721	GAC	ral	ser	len	arg	thr
arg val pro ile	glu vai	. asn pn	e thr	gin	2191/	vai /731	ала	vai	asp	Vul	001			
2161/721 CCC GAC GGC GTC	CCC CTC	י ככייי כא	A CCG	CTC	CGG	ፓጥር	ጥርር	GTG	CAC	TCC	AAC	GCC	TAC	GGC
pro asp gly val	GCG CTG	alval	n ccc	val	ard	leu	ser	val	his	ser	asn	ala	tyr	gly
	ara rec	r gry gr	u pro	Val	2251	/751							-	-
2221/741 AAG GTG TTG TTC	GCG ATO	ACG CT	A TCC	GCT	GCG (GCC	GTG	CTG	GTA	ACG	CTG	GCG	GGC	CGG
lys val leu phe	ala ile	thr le	u ser	ala	ala a	ala	val	leu	val	thr	leu	ala	gly	arg
2201/761					2311.	///1								
CCC CTT TCC CAC	CGG TTC	C CGT GG	C CAG	CCT	GAT (CGC	GCC	GAC	CTG	GAT	CGC	CCC	GAC	CTG
arg leu trp his	arg phe	e arg gl	y gln	pro	asp	arg	ala	asp	leu	asp	arg	pro	asp	leu
2211/781					23/1,	/ /91								
CCT ACC GGC AAA	CAC GC	C CCG CA	G CGC	CGT	GCC	GTA	GCC	AGT	CGG	GAT	' GAC	GAA	AAG	CAC
pro thr gly lys	his ala	a pro gl	n arg	arg	ala '	val	ala	ser	arg	asp	asp	glu	тла	nls
2401/801														
CGG GTA TGA														
arg val OPA														

SEQ ID No.36D (continued 2)

FIGURE 36D (continued 2)



ORF according to Cole et al., 1998 (Nature 393 537-544) and containing Rv 3909.

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31/11
1/1
TGA CTC AGC ACC GGG TCA GCA CAA CGG TCC CGG GCC GGG GCC GTG ACC GCA CTG CAA CTC
OPA leu ser thr gly ser ala gln arg ser arg ala gly ala val thr ala leu gln leu
                                        91/31
GGC TGG GCC GCT TTG GCG CGC GTC ACC TCA GCG ATC GGC GTC GTG GCC GGC CTC GGG ATG
gly trp ala ala leu ala arg val thr ser ala ile gly val val ala gly leu gly met
                                        151/51
GCG CTC ACG GTA CCG TCG GCG GCA CCG CAC GCG CTC GCA GGC GAG CCC AGC CCG ACG CCT
ala leu thr val pro ser ala ala pro his ala leu ala gly glu pro ser pro thr pro
                                        211/71
TTT GTC CAG GTC CGC ATC GAT CAG GTG ACC CCG GAC GTG GTG ACC ACT TCC AGC GAA CCC
phe val gln val arg ile asp gln val thr pro asp val val thr thr ser ser glu pro
                                        271/91
241/81
CAT GTC ACC GTC AGC GGA ACG GTG ACC AAT ACC GGT GAC CGC CCA GTC CGC GAT GTG ATG
his val thr val ser gly thr val thr asn thr gly asp arg pro val arg asp val met
                                         331/111
301/101
GTC CGG CTT GAG CAC GCC GCG GTC ACG TCG TCA ACG GCG TTA CGC ACC TCG CTC GAC
val arg leu glu his ala ala ala val thr ser ser thr ala leu arg thr ser leu asp
                                         391/131
361/121
GGC GGC ACC GAC CAG TAC CAG CCG GCC GCG GAC TTC CTC ACG GTC GCC CCC GAA CTA GAC
gly gly thr asp gln tyr gln pro ala ala asp phe leu thr val ala pro glu leu asp
                                         451/151
421/141
CGC GGG CAA GAG GCC GGC TTT ACC CTC TCG GCC CCG CTG CGC TCG CTG ACC AGG CCG TCG
arg gly gln glu ala gly phe thr leu ser ala pro leu arg ser leu thr arg pro ser
                                         511/171
TTG GCC GTC AAC CAG CCC GGG ATC TAC CCG GTC CTG GTC AAC GTC AAT GGG ACA CCC GAC
leu ala val asn gln pro gly ile tyr pro val leu val asn val asn gly thr pro asp
                                         571/191
541/181
TAC GGT GCG CCT GCG CGC CTC GAC AAT GCG CGG TTC CTG TTG CCC GTG GTC GGA GTG CCA
tyr gly ala pro ala arg leu asp asn ala arg phe leu leu pro val val gly val pro
                                         631/211
CCC GAC CAG GCC ACC GAC TTC GGC TCC GCT GTT GCA CCA GAA ACG ACG GCG CCG GTC TGG
pro asp gln ala thr asp phe gly ser ala val ala pro glu thr thr ala pro val trp
                                         691/231
 661/221
ATC ACC ATG CTG TGG CCG CTG GCC GAC CGG CCC CGG TTG GCC CCC GGG GCA CCC GGT GGC
 ile thr met leu trp pro leu ala asp arg pro arg leu ala pro gly ala pro gly gly
                                         751/251
 ACC GTT CCC GTC CGG CTG GTC GAC GAC GAC CTG GCA AAC TCG CTG GCC AAC GGC GGC CGG
 thr val pro val arg leu val asp asp leu ala asn ser leu ala asn gly gly arg
                                         811/271
 781/261
 CTG GAC ATC CTC CTG TCG GCC GCC GAG TTC GCC ACC AAC CGG GAA GTC GAC CCC GAC GGC
 leu asp ile leu leu ser ala ala glu phe ala thr asn arg glu val asp pro asp gly
                                         871/291
 841/281
 GCC GTC GGC CGA GCG CTG TGC CTG GCC ATC GAC CCA GAT CTA CTC ATC ACC GTC AAT GCG
 ala val gly arg ala leu cys leu ala ile asp pro asp leu leu ile thr val asn ala
                                         931/311
 901/301
 ATG ACC GGC GGC TAC GTC GTC GAC TCG CCC GAC GGG GCC GCT CAA CTA CCG GGC ACC
 met thr gly gly tyr val val ser asp ser pro asp gly ala ala gln leu pro gly thr
                                         991/331
 961/321
 CCG ACC CAC CCG GGC ACC GGC CAG GCC GCA TCC AGC TGG CTG GAT CGA TTG CGG ACG
 pro thr his pro gly thr gly gln ala ala ala ser ser trp leu asp arg leu arg thr
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SEQ ID No.36F

FIGURE 36F



1021/341								1051/									
CTA GTC CAC	CGG	ACA	TGC	GTG	ACG	CCG	CTG	CCT T	TT	GCC	CAA	GCC	GAC	CTG	GAT	GCT	TTG
leu val his	arg	thr	cys	val	thr	pro	leu	pro p	he	ala	gln	ala	asp	leu	asp	ala	leu
1081/361								1111/	'371								
CAG CGG GTT	TAA	GAT	CCG	AGG	CTG	AGC	GCG	ATC G	GCA .	ACC	ATC	AGC	CCC	GCC	GAC	ATC	GTC
gln arg val	asn	asp	pro	arg	leu	ser	ala	ile a	ıla	thr	ile	ser	pro	ala	asp	ile	val
1141/381								1171/	'391								
GAC CGC ATC	CTG	GAT	GTC	AGC	TCC	ACC	CGC	GGC G	SCA.	ACC	GTG	CTG	CCC	GAC	GGC	CCG	TTG
asp arg ile	leu	asp	val	ser	ser	thr	arg	gly a	ala	thr	val	leu	pro	asp	дтй	pro	ıeu
1201/401								1231/			7.00	CITIE	ccc	CTTC	CCG	CCC	ccc
ACC GGC CGG	GCG	ATC	AAC	TTG	CTC	AGC	ACC	CAC G	-1	AAC	+hr	7731	212	Tral	ala	ala	ala
thr gly arg	ala	ile	asn	Leu	leu	ser	tnr	1291/	//31 //31	asii	CIII	vaı	ата	Val	ала	ата	ala
1261/421 GAT TTT AGC	~~~	~ ~ ~	C 7 7	CAC	CAC	CCT	ሞርር				GGC	TCC	GCG	CTC	тта	CCC	GCT
GAT TTT AGC asp phe ser	CCC	GAG	GAA	CAG	CAG aln	414	car	ser (rln	ile	alv	ser	ala	leu	leu	pro	ala
	pro	gru	gru	gin	gin	gry	361	1351	/451		9-1					L	
1321/441 ACC GCG CCC	ccc	ccc	ጥጥር	ጥርር	CCG	CGG	GTG				CCG	TTT	GAT	CCC	GCG	GTC	GGG
thr ala pro	ara	ard	leu	ser	pro	arg	val	val a	ala	ala	pro	phe	asp	pro	ala	val	gly
1381/461								1411,	/471	_							
CCC CCG CTG	GCC	GCC	GCG	GGA	ACA	AAC	CCG	ACC (GTT	CCT	ACC	TAT	CTA	GAT	CCC	TCG	TTG
ala ala leu	ala	ala	ala	gly	thr	asn	pro	thr	val	pro	thr	tyr	leu	asp	pro	ser	leu
1//1//81								1471,	/491	L							
TTC GTT CGG	ATC	GCG	CAT	GAA	TCG	ATC	ACC	GCG (CGC	CGC	CAG	GAC	GCC	TTG	GGC	GCA	ATG
phe val arg	ile	ala	his	glu	ser	ile	thr	ala a	arg	arg	gln	asp	ala	leu	gly	ala	met
1501/501								1531.	/51]	L 	~		am a	cmc	aaa	ccc	ccc
CTG TGG CGC	AGC	TTG	GAG	CCG	AAT	GCC	GCG	CCC (CGT	ACC	CAA	ATC	CTG	GTG	220	220	313
leu trp arg	ser	leu	glu	pro	asn	ala	ala	pro	arg	tnr	grn	ше	Teu	vai	pro	Pro	ала
1561/521 TCG TGG AGC	~~~	~~~	7.00	a	CAC	ccc	CNC	1591			A C C	ccc	CTG	GCC	ACC	GCC	ATC
TCG TGG AGC ser trp ser	CTG	GCC	AGC	GAC	GAC	212	cAG aln	ual	ila	len	thr	ala	leu	ala	thr	ala	ile
	leu	ala	ser	asp	asp	ala	giii	1651	/55°	1 Cu	CIII	ulu	104	414			
1621/541 CGG TCT GGC	CTTC	ccc	CTC	ccc	CGA	CCA	СТА	CCG	GCG	- GTG	ATC	GCT	GAC	GCC	GCG	GCC	CGC
arg ser gly	leu	ala	val	pro	arg	pro	leu	pro	ala	val	ile	ala	asp	ala	ala	ala	arg
1691/561								1711	/57:	1							
ACC GAG CCA	CCG	GAA	CCC	CCG	GGC	GCT	TAC	AGC	GCC	GCT	CGC	GGC	CGG	TTC	AAT	GAC	GAC
thr glu pro	pro	glu	pro	pro	gly	ala	tyr	ser	ala	ala	arg	gly	arg	phe	asn	asp	asp
17/1/591								1771	/59	1							
ATC ACC ACG	CAG	ATC	GGC	GGG	CAG	GTT	GCC	CGG	CTA	TGG	AAG	CTG	ACC	TCG	GCG	TTG	ACC
ile thr thr	gln	ile	gly	gly	gln	val	ala	arg	leu	trp	lys	leu	thr	ser	ala	leu	thr
1801/601								1831	/61	T							,
ATC GAT GAC	CGC	ACC	GGG	CTG	ACC	GGC	GTG	CAG	TAC	ACC	GCA	CCA	LON	320	GAG	SAC	met
ile asp asp	arg	thr	gly	Leu	thr	đΤλ	val	. gin	tyr	1 CUI	ala	pro	Teu	ary	gru	asp	mee
1861/621				шаа	am a	CCI	~~~	1891	./ 63 7.CC	T	ממי	ccc	ርጥር	GCC	CAG	CAG	CGG
TTG CGC GCG leu arg ala	CTG	AGC	CAA	TCG	CTA	CCA		GAT	+hr	2 70	asn	alv	len	ala	aln	aln	arg
	leu	ser	gın	ser	Teu	pro	pro	, asp 1951	/65	1	usn	9-1			9	5	. ,
1921/641 CTG GCC GTC	Cmm	CCA	. אא	. Acc	: ውጥረ	GAC	ር ይ ፕ	י Стт בסב	., 55 ግጥሮ	GGC	GCG	GTG	ACC	ATC	GTC	AAC	CCG
leu ala val	GII	~1 ·	, lue	the	, Alc	asn	asr	leu	phe	alv	ala	val	thr	ile	val	asn	pro
1001/661								2011	_/ 67	1							
CCC CCC TCC	ተልር	ACT	CTG	GCC	: ACC	GAG	CAC	AGT	CCG	CTG	CCG	TTG	GCG	CTG	CAT	'AA'	GGC
gly gly ser	tvr	thr	leu	ala	thr	glu	his	ser	pro	leu	pro	leu	ala	leu	his	asr	gly
9-1 9-1 -01	~ <u>, </u>					~											

SEQ ID 36F (continued 1)

FIGURE 36F (continued 1)



2071/691 2041/681 CTC GCC GTG CCA ATC CGG GTC CGG CTA CAG GTC GAT GCT CCG CCC GGG ATG ACG GTG GCC leu ala val pro ile arg val arg leu gln val asp ala pro pro gly met thr val ala 2131/711 GAT GTC GGT CAG ATC GAG CTA CCG CCC GGG TAC CTG CCG CTA CGA GTA CCA ATC GAG GTG asp val gly gln ile glu leu pro pro gly tyr leu pro leu arg val pro ile glu val 2191/731 2161/721 AAC TTC ACA CAG CGG GTT GCC GTC GAC GTG TCG CTG CGG ACC CCC GAC GGC GTC GCG CTG asn phe thr gln arg val ala val asp val ser leu arg thr pro asp gly val ala leu 2251/751 2221/741 GGT GAA CCG GTG CGG TTG TCG GTG CAC TCC AAC GCC TAC GGC AAG GTG TTG TTC GCG ATC gly glu pro val arg leu ser val his ser asn ala tyr gly lys val leu phe ala ile 2311/771 2281/761 ACG CTA TCC GCT GCG GCC GTG CTG GTA ACG CTG GCG GGC CGG CGC CTT TGG CAC CGG TTC thr leu ser ala ala ala val leu val thr leu ala gly arg arg leu trp his arg phe 2371/791 2341/781 CGT GGC CAG CCT GAT CGC GCC GAC CTG GAT CGC CCC GAC CTG CCT ACC GGC AAA CAC GCC arg gly gln pro asp arg ala asp leu asp arg pro asp leu pro thr gly lys his ala 2431/811 2401/801 CCG CAG CGC CGT GCC GTA GCC AGT CGG GAT GAC GAA AAG CAC CGG GTA TGA pro gln arg arg ala val ala ser arg asp glu lys his arg val OPA

SEQ ID 36F (continued 2)

FIGURE 36F (continued 2)

31/11 1/1 ATC CGC GCG TTG GCG TCG CAT CCG AAC ATC GTC GGA GTC AAG GAC GCC AAA GCC GAC CTG ile arg ala leu ala ser his pro asn ile val gly val lys asp ala lys ala asp leu 91/31 61/21 CAC AGC GGC GCC CAA ATC ATG GCC GAC ACC GGA CTG GCC TAC TAT TCC GGC GAC GAC GCG his ser gly ala gln ile met ala asp thr gly leu ala tyr tyr ser gly asp asp ala 151/51 CTC AAC CTG CCC TGG CTG GCC ATG GGC GCC ACG GGC TTC ATC AGC GTG ATT GCC CAC CTG leu asn leu pro trp leu ala met gly ala thr gly phe ile ser val ile ala his leu 211/71 GCA GCC GGG CAG CTT CGA GAG TTG TTG TCC GCC TTC GGT TCT GGG GAT ATC GCC ACC GCC ala ala gly gln leu arg glu leu leu ser ala phe gly ser gly asp ile ala thr ala 241/81 CGC AAG ATC arg lys ile

SEQ ID No.37A

FIGURE 37A

116/185

31/11 1/1 GAT CCG CGC GTT GGC GTC GCA TCC GAA CAT CGT CGG AGT CAA GGA CGC CAA AGC CGA CCT asp pro arg val gly val ala ser glu his arg arg ser gln gly arg gln ser arg pro 91/31 GCA CAG CGG CGC CCA AAT CAT GGC CGA CAC CGG ACT GGC CTA CTA TTC CGG CGA CGC ala gln arg arg pro asn his gly arg his arg thr gly leu leu phe arg arg arg 151/51 GCT CAA CCT GCC CTG GCT GGC CAT GGG CGC CAC GGG CTT CAT CAG CGT GAT TGC CCA CCT ala gln pro ala leu ala gly his gly arg his gly leu his gln arg asp cys pro pro 211/71 GGC AGC CGG GCA GCT TCG AGA GTT GTT GTC CGC CTT CGG TTC TGG GGA TAT CGC CAC CGC gly ser arg ala ala ser arg val val val arg leu arg phe trp gly tyr arg his arg 241/81 CCG CAA GAT C pro gln asp

SEQ ID No.37B

FIGURE 37B

31/11 1/1 TCC GCG CGT TGG CGT CGC ATC CGA ACA TCG TCG GAG TCA AGG ACG CCA AAG CCG ACC TGC ser ala arg trp arg arg ile arg thr ser ser glu ser arg thr pro lys pro thr cys 91/31 61/21 ACA GCG GCG CCC AAA TCA TGG CCG ACA CCG GAC TGG CCT ACT ATT CCG GCG ACG CGC thr ala ala pro lys ser trp pro thr pro asp trp pro thr ile pro ala thr thr arg 151/51 121/41 TCA ACC TGC CCT GGC TGG CCA TGG GCG CCA CGG GCT TCA TCA GCG TGA TTG CCC ACC TGG ser thr cys pro gly trp pro trp ala pro arg ala ser ser ala OPA leu pro thr trp 211/71 181/61 CAG CCG GGC AGC TTC GAG AGT TGT TGT CCG CCT TCG GTT CTG GGG ATA TCG CCA CCG CCC gln pro gly ser phe glu ser cys cys pro pro ser val leu gly ile ser pro pro 241/81 GCA AGA TC ala arg

SEQ ID No.37C

FIGURE 37C



Coding sequence Rv2753c predicted by Cole et al., 1998 (Nature 393 537-544) containing Seq 37A

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GTG ACC ACC GTC GGA TTC GAC GTC GCA GCG CGC CTA GGA ACC CTG CTG ACC GCG ATG GTG
val thr thr val gly phe asp val ala ala arg leu gly thr leu leu thr ala met val
                                        91/31
61/21
ACA CCG TTT AGC GGC GAT GGC TCC CTG GAC ACC GCC ACC GCG GCG CGG CTG GCC AAC CAC
thr pro phe ser gly asp gly ser leu asp thr ala thr ala ala arg leu ala asn his
                                        151/51
CTG GTC GAT CAG GGG TGC GAC GGT CTG GTG GTC TCG GGC ACC ACC GGC GAG TCG CCG ACC
leu val asp gln gly cys asp gly leu val val ser gly thr thr gly glu ser pro thr
                                        211/71
181/61
ACC ACC GAC GGG GAG AAA ATC GAG CTG CTG CGG GCC GTC TTG GAA GCG GTG GGG GAC CGG
thr thr asp gly glu lys ile glu leu leu arg ala val leu glu ala val gly asp arg
                                        271/91
GCC CGT GTT ATC GCC GGT GCC GGC ACC TAT GAC ACC GCG CAC AGC ATC CGG CTG GCC AAG
ala arg val ile ala gly ala gly thr tyr asp thr ala his ser ile arg leu ala lys
                                        331/111
301/101
GCT TGT GCG GCC GAG GGT GCG CAC GGG CTG CTG GTG GTC ACG CCC TAC TAT TCC AAG CCG
ala cys ala ala glu gly ala his gly leu leu val val thr pro tyr tyr ser lys pro
                                         391/131
361/121
CCG CAG CGG GGG CTG CAA GCC CAT TTC ACC GCC GTC GCC GAC GCG ACC GAG CTG CCG ATG
pro gln arg gly leu gln ala his phe thr ala val ala asp ala thr glu leu pro met
                                         451/151
421/141
CTG CTC TAT GAC ATC CCG GGG CGG TCG GCG GTG CCG ATC GAG CCC GAC ACG ATC CGC GCG
leu leu tyr asp ile pro gly arg ser ala val pro ile glu pro asp thr ile arg ala
                                         511/171
TTG GCG TCG CAT CCG AAC ATC GTC GGA GTC AAG GAC GCC AAA GCC GAC CTG CAC AGC GGC
leu ala ser his pro asn ile val gly val lys asp ala lys ala asp leu his ser gly
                                         571/191
GCC CAA ATC ATG GCC GAC ACC GGA CTG GCC TAC TAT TCC GGC GAC GCG CTC AAC CTG
ala gln ile met ala asp thr gly leu ala tyr tyr ser gly asp asp ala leu asn leu
                                         631/211
CCC TGG CTG GCC ATG GGC GCC ACG GGC TTC ATC AGC GTG ATT GCC CAC CTG GCA GCC GGG
pro trp leu ala met gly ala thr gly phe ile ser val ile ala his leu ala ala gly
                                         691/231
661/221
CAG CTT CGA GAG TTG TCC GCC TTC GGT TCT GGG GAT ATC GCC ACC GCC CGC AAG ATC
gln leu arg glu leu leu ser ala phe gly ser gly asp ile ala thr ala arg lys ile
                                         751/251
721/241
AAC ATT GCG GTC GCC CCG CTG TGC AAC GCG ATG AGC CGC CTG GGT GGG GTG ACG TTG TCC
asn ile ala val ala pro leu cys asn ala met ser arg leu gly gly val thr leu ser
                                         811/271
AAG GCG GGC TTG CGG CTG CAG GGC ATC GAC GTC GGT GAT CCC CGG CTG CCC CAG GTG GCC
lys ala gly leu arg leu gln gly ile asp val gly asp pro arg leu pro gln val ala
                                         871/291
 841/281
GCG ACA CCG GAG CAG ATC GAC GCG TTG GCC GCC GAC ATG CGC GCG GCC TCG GTG CTT CGG
 ala thr pro glu gln ile asp ala leu ala ala asp met arg ala ala ser val leu arg
```

901/301 TGA OPA

SEQ ID No.37D

FIGURE 37D



ORF according to Cole et al., 1998 (Nature 393 537-544) containing Rv2753c

```
31/11
1/1
TAA GGT GAG CGC CGT GGC CGA GAC CGC GCC GCT GCG CGT GCA ACT GAT CGC CAA GAC CGA
OCH gly glu arg arg gly arg asp arg ala ala arg ala thr asp arg gln asp arg
                                       91/31
61/21
CTT CTT GGC CCC ACC CGA CGT GCC CTG GAC CAC CGA CGC CGG CGG ACC CGC GCT GGT
leu leu gly pro thr arg arg ala leu asp his arg arg arg arg thr arg ala gly
                                        151/51
CGA GTT CGC CGG CCG GGC CTG CTA TCA GAG CTG GTC CAA GCC CAA TCC CAA GAC CGC CAC
arg val arg arg pro gly leu leu ser glu leu val gln ala gln ser gln asp arg his
                                        211/71
181/61
CAA CGC CGG CTA CCT CCG GCA CAT CAT CGA CGT CGG ACA TTT CTC GGT GCT AGA GCA TGC
gln arg arg leu pro pro ala his his arg arg thr phe leu gly ala arg ala cys
                                        271/91
CAG CGT GTC GTT CTA CAT CAC CGG GAT CTC GCG ATC GTG CAC CCA CGA GCT GAT CCG CCA
gln arg val val leu his his arg asp leu ala ile val his pro arg ala asp pro pro
                                        331/111
301/101
CCG GCA TTT CTC CTA CTC GCA GCT CTC CCA GCG CTA CGT ACC CGA GAA GGA CTC GCG GGT
pro ala phe leu leu ala ala leu pro ala leu arg thr arg glu gly leu ala gly
                                        391/131
CGT CGT GCC GCC CGG CAT GGA GGA CGC CGA CCT GCG CCA CAT CCT GAC CGA GGC CGC
arg arg ala ala arg his gly gly arg arg pro ala pro his pro asp arg gly arg
                                        451/151
421/141
CGA CGC CGC CGC CAC CTA CAG CGA GCT GCT GGC CAA GCT GGA AGC CAA GTT CGC CGA
arg arg pro arg his leu gln arg ala ala gly gln ala gly ser gln val arg arg
                                        511/171
481/161
CCA ACC CAA CGC GAT CCT GCG CCG CAA GCA GGC CCG CCA AGC CGC CCG CGC GGT GCT GCC
pro thr gln arg asp pro ala pro gln ala gly pro pro ser arg pro arg gly ala ala
                                        571/191
541/181
CAA CGC CAC CGA AAC CCG CAT CGT GGT GAC CGG CAA CTA CCG GGC CTG GCG GCA CTT CAT
gln arg his arg asn pro his arg gly asp arg gln leu pro gly leu ala ala leu his
                                        631/211
601/201
CGC AAT GCG GGC CAG CGA GCA CGC CGA CGT GGA AAT CCG GCG ACT GGC CAT CGA ATG CCT
arg asn ala gly gln arg ala arg arg gly asn pro ala thr gly his arg met pro
                                        691/231
661/221
GCG CCA GCT CGC CGT GGC CCC CGC GGT GTT CGC CGA CTT CGA GGT GAC CAC CCT GGC
ala pro ala arg arg gly pro arg gly val arg arg leu arg gly asp his pro gly
                                        751/251
721/241
CGA CGG CAC CGA GGT GGC CAG CCC GTT GGC GAC CGA AGC CTG AGG CGG CGT GTC GCT
arg arg his arg gly gly asp gln pro val gly asp arg ser leu arg arg arg val ala
                                        811/271
781/261
GGA CAA ACA CGC GCG CTC GCG GCC GGG ATA AAG CGC CAG GTA ACC TTG GGA GCC GTG ACC
gly gln thr arg ala leu ala ala gly ile lys arg gln val thr leu gly ala val thr
                                        871/291
 841/281
ACC GTC GGA TTC GAC GTC GCA GCG CGC CTA GGA ACC CTG CTG ACC GCG ATG GTG ACA CCG
 thr val gly phe asp val ala ala arg leu gly thr leu leu thr ala met val thr pro
                                        931/311
 901/301
 TTT AGC GGC GAT GGC TCC CTG GAC ACC GCC ACC GCG GCG CTG GCC AAC CAC CTG GTC
 phe ser gly asp gly ser leu asp thr ala thr ala ala arg leu ala asn his leu val
                                         991/331
 961/321
 GAT CAG GGG TGC GAC GGT CTG GTG GTC TCG GGC ACC ACC GGC GAG TCG CCG ACC ACC
 asp gln gly cys asp gly leu val val ser gly thr thr gly glu ser pro thr thr thr
```

SEQ ID No.37F

FIGURE 37F



1021/341	1051/351
GAC GGG GAG AAA ATC GAG CTG CTG CGG GCC	GTC TTG GAA GCG GTG GGG GAC CGG GCC CGT
asp gly glu lys ile glu leu leu arg ala	val leu glu ala val gly asp arg ala arg
1081/361	1111/371
GTT ATC GCC GGT GCC GGC ACC TAT GAC ACC	GCG CAC AGC ATC CGG CTG GCC AAG GCT TGT
val ile ala gly ala gly thr tyr asp thr	ala his ser ile arg leu ala lys ala cys
11/1/381	1171/391
CCC CCC CAC GGT GCG CAC GGG CTG CTG GTG	GTC ACG CCC TAC TAT TCC AAG CCG CCG CAG
ala ala glu gly ala his gly leu leu val	val thr pro tyr tyr ser lys pro pro gln
1201/401	1231/411
CGG GGG CTG CAA GCC CAT TTC ACC GCC GTC	GCC GAC GCG ACC GAG CTG CCG ATG CTC
arg gly leu gln ala his phe thr ala val	ala asp ala thr glu leu pro met leu leu
1261/421	1291/431
TAT GAC ATC CCG GGG CGG TCG GCG GTG CCG	ATC GAG CCC GAC ACG ATC CGC GCG TTG GCG
tyr asp ile pro gly arg ser ala val pro	ile glu pro asp thr ile arg ala leu ala
1321/441	1351/451
TCG CAT CCG AAC ATC GTC GGA GTC AAG GAC	GCC AAA GCC GAC CTG CAC AGC GGC GCC CAA
ser his pro asn ile val gly val lys asp	ala lys ala asp leu his ser gly ala gln
1381/461	1411/471
ATC ATG GCC GAC ACC GGA CTG GCC TAC TAT	TCC GGC GAC GAC GCG CTC AAC CTG CCC TGG
	ser gly asp asp ala leu asn leu pro trp
1441/481	1471/491
CTG GCC ATG GGC GCC ACG GGC TTC ATC AGC	GTG ATT GCC CAC CTG GCA GCC GGG CAG CTT
	val ile ala his leu ala ala gly gln leu 1531/511
1501/501	GAT ATC GCC ACC GCC CGC AAG ATC AAC ATT
CGA GAG TTG TTG TCC GCC TTC GGT TCT GGG	asp ile ala thr ala arg lys ile asn ile
arg giu leu leu ser ala phe gly ser gly 1561/521	1591/531
1561/521	CGC CTG GGT GGG GTG ACG TTG TCC AAG GCG
GCG GTC GCC CCG CTG TGC AAC GCG ATG NCC	arg leu gly gly val thr leu ser lys ala
1621/541	1651/551
CCC TTC CCC CTC CAG GGC ATC GAC GTC GGT	GAT CCC CGG CTG CCC CAG GTG GCC GCG ACA
alv leu arg leu glo gly ile asp val gly	asp pro arg leu pro gln val ala ala thr
1681/561	1711/571
CCG GAG CAG ATC GAC GCG TTG GCC GCC GAC	
pro glu gln ile asp ala leu ala ala asp	met arg ala ala ser val leu arg OPA
bro 314 311 110 45h 111 111 111 111 1	

SEQ ID No.37F (continued 1)

FIGURE 37F (continued 1)



31/11 1/1 GCG GTG AAC TGG TGG GCC CGG ATG GTT CAA GTA CGC CGT CGC AAA CTC GAG CAC AAC AGG ala val asn trp trp ala arg met val gln val arg arg arg lys leu glu his asn arg 91/31 61/21AGA CGA CGG ATG GAA GGA GAT GCT GGC GCC GGC CAG CTG AAC CCT GCC GAT GCG AAT AAG arg arg met glu gly asp ala gly ala gly gln leu asn pro ala asp ala asn lys 151/51 121/41 TCG TCG TCT ACG GAG GTG AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GCC GAC CAG ser ser ser thr glu val lys ala ala asp ser ala glu ser asp ala gly ala asp gln 211/71 ACT GGC CCG CAG GTG AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GAG CTC GGC GAG thr gly pro gln val lys ala ala asp ser ala glu ser asp ala gly glu leu gly glu 271/91 GAC GCG TGC CCA GAA CAG GCC CTC GTC GAG CGG CGC CCG TCG CGG TTG CGG CGA GGC TGG asp ala cys pro glu gln ala leu val glu arg arg pro ser arg leu arg arg gly trp 331/111 301/101 CTT GTT GGC ATT GCG GCG ACG CTG CTC GCG TTG GCC GGT GGC CTT GGC GCA GCG GGT TAT leu val gly ile ala ala thr leu leu ala leu ala gly gly leu gly ala ala gly tyr 391/131 361/121 TTT GCG TTG CGC TCA CAC CAG GAA AGC CAA TCA ATC GCG CGC GAG GAC CTT GCG GCC ATT phe ala leu arg ser his gln glu ser gln ser ile ala arg glu asp leu ala ala ile 451/151 GAG GCC GCT AAG GAT TGC GTT GCG GCC ACG CAG GCA CCC GAT GCT GGG GCG ATG TCG GCT glu ala ala lys asp cys val ala ala thr gln ala pro asp ala gly ala met ser ala 481/161 AGC ATG CAG AAG ATC ser met gln lys ile

SEQ ID No.38A

FIGURE 38A

31/11 1/1 CAG CGG TGA ACT GGT GGG CCC GGA TGG TTC AAG TAC GCC GTC GCA AAC TCG AGC ACA ACA gln arg OPA thr gly gly pro gly trp phe lys tyr ala val ala asn ser ser thr thr 91/31 61/21 GGA GAC GAC GGA TGG AAG GAG ATG CTG GCG CCG GCC AGC TGA ACC CTG CCG ATG CGA ATA gly asp asp gly trp lys glu met leu ala pro ala ser OPA thr leu pro met arg ile 151/51 121/41 AGT CGT CGT CTA CGG AGG TGA AGG CGG CGG ATT CGG CGG AAT CTG ACG CCG GAG CCG ACC ser arg arg leu arg arg OPA arg arg ile arg arg asn leu thr pro glu pro thr 211/71 181/61 AGA CTG GCC CGC AGG TGA AGG CGG CGG ATT CGG CGG AAT CTG ACG CCG GAG AGC TCG GCG arg leu ala arg arg OPA arg arg ile arg arg asn leu thr pro glu ser ser ala 271/91 241/81 AGG ACG CGT GCC CAG AAC AGG CCC TCG TCG AGC GGC GCC CGT CGC GGT TGC GGC GAG GCT arg thr arg ala gln asn arg pro ser ser ser gly ala arg arg gly cys gly glu ala 331/111 301/101 GGC TTG TTG GCA TTG CGG CGA CGC TGC TCG CGT TGG CCG GTG GCC TTG GCG CAG CGG GTT gly leu leu ala leu arg arg arg cys ser arg trp pro val ala leu ala gln arg val 391/131 361/121 ATT TTG CGT TGC GCT CAC ACC AGG AAA GCC AAT CAA TCG CGC GCG AGG ACC TTG CGG CCA ile leu arg cys ala his thr arg lys ala asn gln ser arg ala arg thr leu arg pro 451/151 TTG AGG CCG CTA AGG ATT GCG TTG CGG CCA CGC AGG CAC CCG ATG CTG GGG CGA TGT CGG leu arg pro leu arg ile ala leu arg pro arg arg his pro met leu gly arg cys arg 481/161 CTA GCA TGC AGA AGA TC leu ala cys arg arg

SEQ ID No.38B

FIGURE 38B



1/1										31/1									
AGC 0	GT	GAA	CTG	GTG	GGC	CCG	GAT	GGT	TCA	AGT	ACG	CCG	TCG	CAA	ACT	CGA	GCA	CAA	CAG
ser o	ılv	glu	leu	val	gly	pro	asp	gly	ser	ser	thr	pro	ser	gln	thr	arg	ala	gln	gln
61/21	1	_								91/3	31								
GAG A	ACG	ACG	GAT	GGA	AGG	AGA	TGC	TGG	CGC	CGG	CCA	GCT	GAA	CCC	TGC	CGA	TGC	GAA	TAA
glu t	thr	thr	asp	gly	arg	arg	cys	trp	arg	arg	pro	ala	glu	pro	cys	arg	cys	glu	OCH
121/4	4 1									151,	/51								
GTC (GTC	GTC	TAC	GGA	GGT	GAA	GGC	GGC	GGA	TTC	GGC	GGA	ATC	TGA	CGC	CGG	AGC	CGA	CCA
val v	val	val	tyr	gly	gly	glu	gly	gly	gly	phe	gly	gly	ile	OPA	arg	arg	ser	arg	pro
181/	61									211,									
GAC :	rgg	CCC	GCA	GGT	GAA	GGC	GGC	GGA	TTC	GGC	GGA	ATC	TGA	CGC	CGG	AGA	GCT	CGG	CGA
asp 1	trp	pro	ala	gly	glu	gly	gly	gly	phe	gly	gly	ile	OPA	arg	arg	arg	ala	arg	arg
241/8	81									271.								3.00	ome
GGA (CGC	GTG	CCC	AGA	ACA	GGC	CCT	CGT	CGA	GCG	GCG	CCC	GTC	GCG	GTT	GCG	GCG	AGG	CTG
gly	arg	val	pro	arg	thr	gly	pro	arg	arg	ala	ala /111	pro	vaı	ala	vai	ala	ата	ary	Tea
301/	101							a am	~~~		/111	ccc	mcc	CCE	TICC	ccc	λGC	ccc	מיחים
GCT '	TGT	TGG	CAT	TGC	GGC	GAC	GCT	GCT	CGC	GTT	GGC	CGG	TGG +xn	201	+ rn	250	car	alv	len
ala		trp	his	cys	₫Ţλ	asp	aıa	ala	arg	Val	91y /131	ary	стр	PLO	crb	arg	301	9±3	ıcu
361/	121							1 T C	007			ccc	ccc	CCA	CCN	CCT.	TICC	GGC	СУП
TTT '	TGC	GTT	GCG	CTC	ACA	CCA	GGA	AAG	CUA	ATC	AAI	250	212	CGM	alu	nro	CVS	alv	his
phe		val	ala	Leu	thr	pro	дтЛ	туs	pro	151	/151	ary	ата	ary	9±3	pro	Cys	9-1	11110
421/ TGA	141	~~~	CT 70 70	~~1	mmc	CCT	mcc	ccc	CAC	CCA	GGC	ΔCC	CGA	TGC	TGG	GGC	GAT	GTC	GGC
TGA OPA	GGC	CGC	TAA	GGA	116	CGI	160	~1··	hic	ala	~1v	thr	ara	CVS	trn	alv	asp	val	alv
	_	arg	OCH	дтА	reu	arg	Cys	дту	11125	ата	9 + Y	CIII	ary	Cyb	CIP	9-1	u D P		9-1
481/			(1 R R	~ n m	C														
TAG																			
AMB	nıs	aia	$q \perp u$	asp															

SEQ ID No.38C

FIGURE 38C



122/185

Sequence Rv0175 predicted by Cole et al., 1998 (Nature 393 537-544) and containing seq38A

```
31/11
1/1
GTG AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GCC GAC CAG ACT GGC CCG CAG GTG
val lys ala ala asp ser ala glu ser asp ala gly ala asp gln thr gly pro gln val
                                        91/31
AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GAG CTC GGC GAG GAC GCG TGC CCA GAA
lys ala ala asp ser ala glu ser asp ala gly glu leu gly glu asp ala cys pro glu
                                        151/51
CAG GCC CTC GTC GAG CGG CGC CCG TCG CGG TTG CGG CGA GGC TGG CTT GTT GGC ATT GCG
gln ala leu val glu arg arg pro ser arg leu arg arg gly trp leu val gly ile ala
                                        211/71
GCG ACG CTG CTC GCG TTG GCC GGT GGC CTT GGC GCA GCG GGT TAT TTT GCG TTG CGC TCA
ala thr leu leu ala leu ala gly gly leu gly ala ala gly tyr phe ala leu arg ser
                                        271/91
CAC CAG GAA AGC CAA TCA ATC GCG CGC GAG GAC CTT GCG GCC ATT GAG GCC GCT AAG GAT
his gln glu ser gln ser ile ala arg glu asp leu ala ala ile glu ala ala lys asp
                                         331/111
301/101
TGC GTT GCG GCC ACG CAG GCA CCC GAT GCT GGG GCG ATG TCG GCT AGC ATG CAG AAG ATC
cys val ala ala thr gln ala pro asp ala gly ala met ser ala ser met gln lys ile
                                         391/131
361/121
ATC GAG TGT GGC ACC GGT GAT TTC GGT GCC CAG GCG TCG TTG TAC ACC AGC ATG CTC GTC
ile glu cys gly thr gly asp phe gly ala gln ala ser leu tyr thr ser met leu val
                                         451/151
421/141
GAG GCG TAT CAA GCG GCC AGC GTC CAC GTG CAA GTG ACC GAT ATG CGC GCG GCG GTC GAG
glu ala tyr gln ala ala ser val his val gln val thr asp met arg ala ala val glu
                                         511/171
481/161
CGC AAC AAC AAT GAC GGG TCG GTC GAT GTT CTG GTG GCG CTC CGG GTC AAG GTG TCC AAC
arg asn asn asp gly ser val asp val leu val ala leu arg val lys val ser asn
                                         571/191
541/181
ACC GAC TCG GAT GCC CAT GAA GTC GGC TAC CGT CTT CGG GTC CGG ATG GCA CTG GAT GAG
thr asp ser asp ala his glu val gly tyr arg leu arg val arg met ala leu asp glu
                                         631/211
601/201
GGC CGC TAT AAG ATC GCC AAA CTC GAC CAG GTG ACG AAG TGA
gly arg tyr lys ile ala lys leu asp gln val thr lys OPA
```

SEQ ID No.38D

FIGURE 38D



ORF according to Cole et al., 1998 (Nature 393 537-544) containing Rv0175

```
31/11
1/1
TGA ACT GGT GGG GCC GGA TGG TGT CAA GTA CGC CGT CGC AAA CTC GAG CAC AAC AGG AGA
OPA thr gly gly ala gly trp cys gln val arg arg lys leu glu his asn arg arg
                                        91/31
CGA CGG ATG GAA GGA GAT GCT GGC GCC GGC CAG CTG AAC CCT GCC GAT GCG AAT AAG TCG
arg arg met glu gly asp ala gly ala gly gln leu asn pro ala asp ala asn lys ser
                                        151/51
121/41
TCG TCT ACG GAG GTG AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GCC GAC CAG ACT
ser ser thr glu val lys ala ala asp ser ala glu ser asp ala gly ala asp gln thr
                                        211/71
GGC CCG CAG GTG AAG GCG GCG GAT TCG GCG GAA TCT GAC GCC GGA GAG CTC GGC GAG GAC
gly pro gln val lys ala ala asp ser ala glu ser asp ala gly glu leu gly glu asp
                                        271/91
241/81
GCG TGC CCA GAA CAG GCC CTC GTC GAG CGG CGC CCG TCG CGG TTG CGG CGA GGC TGG CTT
ala cys pro glu gln ala leu val glu arg arg pro ser arg leu arg arg gly trp leu
                                         331/111
301/101
GTT GGC ATT GCG GCG ACG CTG CTC GCG TTG GCC GGT GGC CTT GGC GCA GCG GGT TAT TTT
val gly ile ala ala thr leu leu ala leu ala gly gly leu gly ala ala gly tyr phe
                                         391/131
361/121
GCG TTG CGC TCA CAC CAG GAA AGC CAA TCA ATC GCG CGC GAG GAC CTT GCG GCC ATT GAG
ala leu arg ser his gln glu ser gln ser ile ala arg glu asp leu ala ala ile glu
                                         451/151
421/141
GCC GCT AAG GAT TGC GTT GCG GCC ACG CAG GCA CCC GAT GCT GGG GCG ATG TCG GCT AGC
ala ala lys asp cys val ala ala thr gln ala pro asp ala gly ala met ser ala ser
                                         511/171
481/161
ATG CAG AAG ATC ATC GAG TGT GGC ACC GGT GAT TTC GGT GCC CAG GCG TCG TTG TAC ACC
met gln lys ile ile glu cys gly thr gly asp phe gly ala gln ala ser leu tyr thr
                                         571/191
541/181
AGC ATG CTC GTC GAG GCG TAT CAA GCG GCC AGC GTC CAC GTG CAA GTG ACC GAT ATG CGC
ser met leu val glu ala tyr gln ala ala ser val his val gln val thr asp met arg
                                         631/211
GCG GCG GTC GAG CGC AAC AAC AAT GAC GGG TCG GTC GAT GTT CTG GTG GCG CTC CGG GTC
ala ala val glu arg asn asn asp gly ser val asp val leu val ala leu arg val
                                         691/231
661/221
AAG GTG TCC AAC ACC GAC TCG GAT GCC CAT GAA GTC GGC TAC CGT CTT CGG GTC CGG ATG
lys val ser asn thr asp ser asp ala his glu val gly tyr arg leu arg val arg met
                                         751/251
721/241
GCA CTG GAT GAG GGC CGC TAT AAG ATC GCC AAA CTC GAC CAG GTG ACG AAG TGA
ala leu asp glu gly arg tyr lys ile ala lys leu asp gln val thr lys OPA
```

SEQ ID No.38F

FIGURE 38F



SEQ ID No.39A

FIGURE 39A

1/1
CAC CTC CCC CCC CGC CGC CGC TGC CGC CGG TTC CCT TTC CCA AGG AAT GTC CGG CGC CGG his leu pro pro arg arg arg cys arg arg phe pro phe pro arg asn val arg arg arg 61/21
GCG TGA TGC AAG GCT GCC TTG AGA GCA CCA GCG GCT TGA TCA TGG GCA TCG ACA GCA AGA ala OPA cys lys ala ala leu arg ala pro ala ala OPA ser trp ala ser thr ala arg 121/41
CCG CAC TGG TCG CCG AGC GCA TCA CCG GTG CCG TCG AGG AGA TC
pro his trp ser pro ser ala ser pro val pro ser arg arg

SEQ ID No.39B

FIGURE 39B

1/1
GAC ACC TCC CCC CCC GCC GCC GCT GCC GCC GGT TCC CTT TCC CAA GGA ATG TCC GGC GCC asp thr ser pro pro ala ala ala ala ala gly ser leu ser gln gly met ser gly ala 91/31
GGG CGT GAT GCA AGG CTG CCT TGA GAG CAC CAG CGG CTT GAT CAT GGG CAT CGA CAG CAA gly arg asp ala arg leu pro OPA glu his gln arg leu asp his gly his arg gln gln 121/41
GAC CGC ACT GGT CGC CGA GCG CAT CAC CGG TGC CGT CGA GAG GAT C asp arg thr gly arg arg ala his his arg cys arg arg gly asp

SEQ ID No.39C

FIGURE 39C





125/185

Coding sequence Rv3006 predicted by Cole et al., 1998 (Nature 393 537-544) and containing seq39A

```
1/1
                                         31/11
ATG TGG ACA ACG CGG TTG GTT CGA TCC GGA CTC GCC GCG CTG TGC GCG GCA GTG CTG GTA
Met trp thr thr arg leu val arg ser gly leu ala ala leu cys ala ala val leu val
• 61/21
                                         91/31
 TCG AGC GGC TGC GCA CGG TTC AAC GAC GCT CAA TCT CAG CCG TTC ACC ACC GAA CCG GAG
 ser ser gly cys ala arg phe asn asp ala gln ser gln pro phe thr thr glu pro glu
                                         151/51
 121/41
 CTG CGG CCC CAA CCC AGC TCG ACA CCT CCC CCC CCG CCG CCG CTG CCG CTG CCC GTT CCC TTT
 leu arg pro gln pro ser ser thr pro pro pro pro pro leu pro pro val pro phe
 181/61
                                         211/71
 CCC AAG GAA TGT CCG GCG CCG GGC GTG ATG CAA GGC TGC CTT GAG AGC ACC AGC GGC TTG
pro lys glu cys pro ala pro gly val met gln gly cys leu glu ser thr ser gly leu
                                         271/91
 241/81
ATC ATG GGC ATC GAC AGC AAG ACC GCA CTG GTC GCC GAG CGC ATC ACC GGT GCC GTC GAG
 ile met gly ile asp ser lys thr ala leu val ala glu arg ile thr gly ala val glu
 301/101
                                         331/111
 GAG ATC TCT ATC AGC GCC GAG CCG AAG GTA AAG ACG GTC ATC CCC GTG GAT CCT GCC GGT
 glu ile ser ile ser ala glu pro lys val lys thr val ile pro val asp pro ala gly
                                         391/131
 361/121
 GAC GGT GGC TTG ATG GAC ATT GTG CTG TCG CCC ACC TAC TCG CAA GAC CGG CTG ATG TAC
 asp gly gly leu met asp ile val leu ser pro thr tyr ser gln asp arg leu met tyr
                                         451/151
 421/141
 GCC TAC ATC AGC ACG CCC ACC GAC AAC CGG GTG GTG CGA GTG GCC GAC GGC GAC ATC CCC
 ala tyr ile ser thr pro thr asp asn arg val val arg val ala asp gly asp ile pro
 481/161
                                         511/171
 AAG GAC ATC CTG ACC GGC ATC CCC AAA GGT GCT GCC GGT AAC ACC GGG GCG CTG ATC TTC
 lys asp ile leu thr gly ile pro lys gly ala ala gly asn thr gly ala leu ile phe
 541/181
                                         571/191
 ACC AGT CCC ACC ACG CTG GTC GTG ATG ACC GGG GAT GCT GGC GAC CCG GCG TTG GCC GCC
 thr ser pro thr thr leu val val met thr gly asp ala gly asp pro ala leu ala ala
 601/201
                                         631/211
 GAT CCC CAA TCG TTG GCC GGT AAG GTC CTG CGT ATC GAA CAG CCC ACC ACC ATC GGC CAG
 asp pro gln ser leu ala gly lys val leu arg ile glu gln pro thr thr ile gly gln
                                         691/231
 661/221
ACG CCG CCG ACG ACG GCG CTG TCT GGC ATC GGC TCC GGC GGC GGC TTG TGC ATC GAT CCG
 thr pro pro thr thr ala leu ser gly ile gly ser gly gly gly leu cys ile asp pro
 721/241
                                         751/251
 GTC GAC GGC TCG CTA TAT GTC GCC GAC CGC ACG CCA ACG GCG GAC CGA TTG CAG CGC ATC
 val asp gly ser leu tyr val ala asp arg thr pro thr ala asp arg leu gln arg ile
                                         811/271
 ACC AAG AAC TCG GAG GTC TCT ACG GTA TGG ACC TGG CCG GAC AAG CCC GGC GTG GCC GGG
 thr lys asn ser glu val ser thr val trp thr trp pro asp lys pro gly val ala gly
 841/281
                                         871/291
 TGT GCC GCG ATG GAC GGC ACC GTG CTG GTC AAC CTG ATT AAT ACC AAA CTG ACG GTG GCG
 cys ala ala met asp gly thr val leu val asn leu ile asn thr lys leu thr val ala
                                         931/311
 901/301
 GTC CGG CTC GCG CCG TCG ACC GGT GCG GTC ACC GGA GAA CCC GAC GTT GTC CGC AAA GAC
 val arg leu ala pro ser thr gly ala val thr gly glu pro asp val val arg lys asp
                                         991/331
 961/321
 ACT CAT GCG CAT GCG TGG GCA TTA CGG ATG TCG CCG GAC GGC AAC GTC TGG GGA GCC ACC
 thr his ala his ala trp ala leu arg met ser pro asp gly asn val trp gly ala thr
                                         1051/351
 GTC AAC AAG ACC GCC GGC GAC GCC GAG AAG CTC GAC GAT GTG GTG TTC CCG CTG TTC CCG
 val asn lys thr ala gly asp ala glu lys leu asp asp val val phe pro leu phe pro
                                         1111/371
 1081/361
 CAG GGT GGC GGC TTC CCG CGC AAC AAC GAC GAC AAG ACC TGA
 gln gly gly phe pro arg asn asp asp lys thr OPA
```

SEQ ID No.39D

FIGURE 39D



ORF according to Cole et al., 1998 (Nature 393 537-544) and containing Rv3006

```
31/11
1/1
TAA GGC CAT TTA GTG CCG AAT TGG GGA TTT GAG CGG CGC TTT CGC CAG ACA ATC CGC ACA
OCH gly his leu val pro asn trp gly phe glu arg arg phe arg gln thr ile arg thr
                                        91/31
TTG ACC CTG ACC AGC CCA AAA GGC CCC AAT TGG GCC GCC ATG CCG ACA GTG CGC ACC
leu thr leu thr ser pro pro lys gly pro asn trp ala ala met pro thr val arg thr
                                        151/51
121/41
CCG GCA GGT GGC GGC GAT GCC CAC AAT GTC CGT AGC CTG TCG GTC ATG TGG ACA ACG CGG
pro ala gly gly gly asp ala his asn val arg ser leu ser val met trp thr thr arg
                                        211/71
181/61
TTG GTT CGA TCC GGA CTC GCC GCG CTG TGC GCG GCA GTG CTG GTA TCG AGC GGC TGC GCA
leu val arg ser gly leu ala ala leu cys ala ala val leu val ser ser gly cys ala
                                        271/91
241/81
CGG TTC AAC GAC GCT CAA TCT CAG CCG TTC ACC ACC GAA CCG GAG CTG CGG CCC CAA CCC
arg phe asn asp ala gln ser gln pro phe thr thr glu pro glu leu arg pro gln pro
                                        331/111
AGC TCG ACA CCT CCC CCC CCG CCG CTG CCG CCG GTT CCC TTT CCC AAG GAA TGT CCG
ser ser thr pro pro pro pro pro leu pro pro val pro phe pro lys glu cys pro
                                        391/131
361/121
GCG CCG GGC GTG ATG CAA GGC TGC CTT GAG AGC ACC AGC GGC TTG ATC ATG GGC ATC GAC
ala pro gly val met gln gly cys leu glu ser thr ser gly leu ile met gly ile asp
                                        451/151
421/141
AGC AAG ACC GCA CTG GTC GCC GAG CGC ATC ACC GGT GCC GTC GAG GAG ATC TCT ATC AGC
ser lys thr ala leu val ala glu arg ile thr gly ala val glu glu ile ser ile ser
                                        511/171
GCC GAG CCG AAG GTA AAG ACG GTC ATC CCC GTG GAT CCT GCC GGT GAC GGT GGC TTG ATG
ala glu pro lys val lys thr val ile pro val asp pro ala gly asp gly gly leu met
                                        571/191
541/181
GAC ATT GTG CTG TCG CCC ACC TAC TCG CAA GAC CGG CTG ATG TAC GCC TAC ATC AGC ACG
asp ile val leu ser pro thr tyr ser gln asp arg leu met tyr ala tyr ile ser thr
                                         631/211
601/201
CCC ACC GAC AAC CGG GTG GTG CGA GTG GCC GAC GGC GAC ATC CCC AAG GAC ATC CTG ACC
pro thr asp asn arg val val arg val ala asp gly asp ile pro lys asp ile leu thr
                                         691/231
661/221
GGC ATC CCC AAA GGT GCT GCC GGT AAC ACC GGG GCG CTG ATC TTC ACC AGT CCC ACC ACG
gly ile pro lys gly ala ala gly asn thr gly ala leu ile phe thr ser pro thr thr
                                         751/251
CTG GTC GTG ATG ACC GGG GAT GCT GGC GAC CCG GCG TTG GCC GAT CCC CAA TCG TTG
leu val val met thr gly asp ala gly asp pro ala leu ala ala asp pro gln ser leu
                                         811/271
781/261
GCC GGT AAG GTC CTG CGT ATC GAA CAG CCC ACC ACC ATC GGC CAG ACG CCG CCG ACG
ala gly lys val leu arg ile glu gln pro thr thr ile gly gln thr pro pro thr thr
                                         871/291
GCG CTG TCT GGC ATC GGC TCC GGC GGC GGC TTG TGC ATC GAT CCG GTC GAC GGC TCG CTA
 ala leu ser gly ile gly ser gly gly gly leu cys ile asp pro val asp gly ser leu
                                         931/311
 901/301
TAT GTC GCC GAC CGC ACG CCA ACG GCG GAC CGA TTG CAG CGC ATC ACC AAG AAC TCG GAG
 tyr val ala asp arg thr pro thr ala asp arg leu gln arg ile thr lys asn ser glu
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SEQ ID No.39F

FIGURE 39F



991/331 961/321 GTC TCT ACG GTA TGG ACC TGG CCG GAC AAG CCC GGC GTG GCC GGG TGT GCC GCG ATG GAC val ser thr val trp thr trp pro asp lys pro gly val ala gly cys ala ala met asp 1051/351 GGC ACC GTG CTG GTC AAC CTG ATT AAT ACC AAA CTG ACG GTG GCG GTC CGG CTC GCG gly thr val leu val asn leu ile asn thr lys leu thr val ala val arg leu ala pro 1111/371 1081/361 TCG ACC GGT GCG GTC ACC GGA GAA CCC GAC GTT GTC CGC AAA GAC ACT CAT GCG CAT GCG ser thr gly ala val thr gly glu pro asp val val arg lys asp thr his ala his ala 1171/391 TGG GCA TTA CGG ATG TCG CCG GAC GGC AAC GTC TGG GGA GCC ACC GTC AAC AAG ACC GCC trp ala leu arg met ser pro asp gly asn val trp gly ala thr val asn lys thr ala 1231/411 1201/401 GGC GAC GCC GAG AAG CTC GAC GAT GTG GTG TTC CCG CTG TTC CCG CAG GGT GGC GGC TTC gly asp ala glu lys leu asp asp val val phe pro leu phe pro gln gly gly phe 1261/421 CCG CGC AAC AAC GAC GAC AAG ACC TGA pro arg asn asn asp asp lys thr OPA

SEQ ID No.39F (continued)

FIGURE 39F (continued)

1/1									31/1	.1								
CAA GGC	Стт	GTT	GAG	CCG	GCG	CAC	GAA	AAC	GAT	CGT	TGT	GTG	TAC	ATT	GGT	GTG	TAT	GGC
glu gly	leu	val	alu	pro	ala	his	glu	asn	asp	arg	cys	val	tyr	ile	gly	val	tyr	gly
61/21									91/3	31								
ጥሮሬ ሬጥጥ	GAA	CGT	GTA	TGT	GCC	CGA	CGA	ATT	GGC	GGA	GCG	CGC	CAG	GGC	GCG	GGG	CTT	GAA
ser val	glu	arg	val	cys	ala	arg	arg	ile	gly	gly	ala	arg	gln	gly	ala	glà	leu	gLu
121/41									151,	/51								
CGT CTC	GGC	GCT	GAC	TCA	GGC	CGC	GAT	CAG	TGC	CGA	GTT	GGA	GAA	CTC	CGC	AAC	CGA	TGC
arg leu	gly	ala	asp	ser	gly	arg	asp	gln	cys	arg	val	gly	glu	leu	arg	asn	arg	cys
181/61									211.	/71								
GTG GCT	TGA	GGG	GTT	GGA	ACC	CAG	AAG	CAC	CGG	CGC	TCG	GCA	TGA	TGA	CGT	GCT	GGG	TGC
val ala	OPA	gly	val	gly	thr	gln	lys	his	arg	arg	ser	ala	OPA	OPA	arg	ala	дŢУ	cys
2/1/81									271	/91								
CAT CGA	TGC	CGC	TCG	CGA	TGA	GTT	CGA	AGC	GTG	AGA	GCA	TCG	CCC	ACT	TCG	CCG	CCG	GAG
asp arg	cys	arg	ser	arg	OPA	val	arg	ser	val	arg	ala	ser	pro	thr	ser	pro	pro	glu
301/101	-								331	/111								
CAG GTG	GTC	GTC	GAC	GCG	AGT	GCC	ATG	GTG	GAT	С								
gln val	val	val	asp	ala	ser	ala	met	val	asp									

SEQ ID No.40A

FIGURE 40A



31/11 1/1 AAG GCC TTG TTG AGC CGG CGC ACG AAA ACG ATC GTT GTG TGT ACA TTG GTG TGT ATG GCT lys ala leu leu ser arg arg thr lys thr ile val val cys thr leu val cys met ala 91/31 CGG TTG AAC GTG TAT GTG CCC GAC GAA TTG GCG GAG CGC GCC AGG GCG CGG GGC TTG AAC arg leu asn val tyr val pro asp glu leu ala glu arg ala arg gly leu asn 151/51 GTC TCG GCG CTG ACT CAG GCC GCG ATC AGT GCC GAG TTG GAG AAC TCC GCA ACC GAT GCG val ser ala leu thr gln ala ala ile ser ala glu leu glu asn ser ala thr asp ala 211/71 TGG CTT GAG GGG TTG GAA CCC AGA AGC ACC GGC GCT CGG CAT GAT GAC GTG CTG GGT GCG trp leu glu gly leu glu pro arg ser thr gly ala arg his asp asp val leu gly ala 271/91 ATC GAT GCC GCT CGC GAT GAG TTC GAA GCG TGA GAG CAT CGC CCA CTT CGC CGC CGG AGC ile asp ala ala arg asp glu phe glu ala OPA glu his arg pro leu arg arg arg ser 331/111 301/101 AGG TGG TCG ACG CGA GTG CCA TGG TGG ATC arg trp ser ser thr arg val pro trp trp ile

SEO ID No.40B

FIGURE 40B

31/11 1/1 AGG CCT TGT TGA GCC GGC GCA CGA AAA CGA TCG TTG TGT GTA CAT TGG TGT GTA TGG CTC arg pro cys OPA ala gly ala arg lys arg ser leu cys val his trp cys val trp leu 91/31 61/21 GGT TGA ACG TGT ATG TGC CCG ACG AAT TGG CGG AGC GCG CCA GGG CGC GGG GCT TGA ACG gly OPA thr cys met cys pro thr asn trp arg ser ala pro gly arg gly ala OPA thr 151/51 121/41 TCT CGG CGC TGA CTC AGG CCG CGA TCA GTG CCG AGT TGG AGA ACT CCG CAA CCG ATG CGT ser arg arg OPA leu arg pro arg ser val pro ser trp arg thr pro gln pro met arg 211/71 181/61 GGC TTG AGG GGT TGG AAC CCA GAA GCA CCG GCG CTC GGC ATG ATG ACG TGC TGG GTG CGA gly leu arg gly trp asn pro glu ala pro ala leu gly met met thr cys trp val arg 271/91 TCG ATG CCG CTC GCG ATG AGT TCG AAG CGT GAG AGC ATC GCC CAC TTC GCC GCA GCA ser met pro leu ala met ser ser lys arg glu ser ile ala his phe ala ala gly ala 301/101 GGT GGT CGA CGC GAG TGC CAT GGT GGA TC gly gly arg arg glu cys his gly gly

SEQ ID No.40C

FIGURE 40C



Coding sequence Rv0549c predicted by Cole et al., 1998 (Nature 393:537-544) and containing seq40A

31/11 gtg aga gca tcg ccc act tcg ccg ccg gag cag gtg gtc gtc gac gcg agt gcc atg gtg val arg ala ser pro thr ser pro pro glu gln val val asp ala ser ala met val 91/31 61/21 gat cta ctg gct cgc act agc gat cgg tgc tct gcg gtg cgc gcg cgg ctg gct cgg acc asp leu leu ala arg thr ser asp arg cys ser ala val arg ala arg leu ala arg thr 151/51 gcg atg cac gcg ccg gcg cac ttc gat gca gag gtg ttg tcg gcg ctg ggg cgc atg cag ala met his ala pro ala his phe asp ala glu val leu ser ala leu gly arg met gln 211/71 181/61 cgc gcc ggc gca ctc acc gtt gcc tat gtc gat gcg gca ctg gag gag ttg cga cag gtg arg ala gly ala leu thr val ala tyr val asp ala ala leu glu glu leu arg gln val 271/91 241/81 ceg gtg act cga cac ggt ctt tcg tcg ctg ctt gct gga gcg tgg tcg cgc cgc gac acc pro val thr arg his gly leu ser ser leu leu ala gly ala trp ser arg arg asp thr 331/111 301/101 ctc cgc ctg acc gat gcc ctc tac gtc gag ctg gcc gaa acg gca ggt ctg gtg ttg ttg leu arg leu thr asp ala leu tyr val glu leu ala glu thr ala gly leu val leu leu 391/131 361/121 acc acc gac gaa aga ttg gca cgc gcc tgg ccc tcg gct cac gcc atc ggc tga thr thr asp glu arg leu ala arg ala trp pro ser ala his ala ile gly OPA

SEQ ID No.40D

FIGURE 40D

ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv0549c

```
31/11
1/1
tga gtt cga agc gtg aga gca tcg ccc act tcg ccg ccg gag cag gtg gtc gtc gac gcg
OPA val arg ser val arg ala ser pro thr ser pro pro glu gln val val asp ala
                                        91/31
agt gcc atg gtg gat cta ctg gct cgc act agc gat cgg tgc tct gcg gtg cgc gcg
ser ala met val asp leu leu ala arg thr ser asp arg cys ser ala val arg ala arg
                                        151/51
121/41
ctg gct cgg acc gcg atg cac gcg ccg gcg cac ttc gat gca gag gtg ttg tcg gcg ctg
leu ala arg thr ala met his ala pro ala his phe asp ala glu val leu ser ala leu
                                        211/71
ggg cgc atg cag cgc gcc gcc gca ctc acc gtt gcc tat gtc gat gcg gca ctg gag gag
gly arg met gln arg ala gly ala leu thr val ala tyr val asp ala ala leu glu glu
                                        271/91
241/81
ttg cga cag gtg ccg gtg act cga cac ggt ctt tcg tcg ctg ctt gct gga gcg tgg tcg
leu arg gln val pro val thr arg his gly leu ser ser leu leu ala gly ala trp ser
                                        331/111
ege ege gae ace ete ege etg ace gat gee ete tae gte gag etg gee gaa acg gea ggt
arg arg asp thr leu arg leu thr asp ala leu tyr val glu leu ala glu thr ala gly
                                        391/131
361/121
ctg gtg ttg ttg acc acc gac gaa aga ttg gca cgc gcc tgg ccc tcg gct cac gcc atc
leu val leu leu thr thr asp glu arg leu ala arg ala trp pro ser ala his ala ile
421/141
ggc tga
gly OPA
```

SEO ID No.40F

FIGURE 40F REPLACEMENT SHEET (RULE 26)



31/11 1/1 CCT GGC CGG GAC GCC TAC GTG TAG CCC GCG GCT AGC ACA GGA TAG CCA TTG TTG TGC GGT pro gly arg asp ala tyr val AMB pro ala ala ser thr gly AMB pro leu leu cys gly 91/31 AGC GCC AAA ACG ATC AGC CCT TCG CGG ACA TGT CAG CAC CCG CCT TGG CCG GGA GAG CGG ser ala lys thr ile ser pro ser arg thr cys gln his pro pro trp pro gly glu arg 151/51 CGT CGT GAC CGT GCT GTC ACC ACG TCT GGT TAG GCT CGG GGC GCG GGC TGG CGC GGA GGA arg arg asp arg ala val thr thr ser gly AMB ala arg gly ala gly trp arg gly gly 211/71 GGT GTG TTG CGG AGG AGG TGT GTT GTA GTG GGG ACG GCG GAT CGG CCG TTG GAC GCC TCG gly val leu arg arg cys val val val gly thr ala asp arg pro leu asp ala ser 271/91 241/81 GCC TTG CGG GAC TGG GCA CAC GCC GTC GTC AGC GAT C ala leu arg asp trp ala his ala val val ser asp

SEQ ID No.41A

FIGURE 41A

1/1								31/1									
כיים ככר ה	GG ACG	CCT	ACG	TGT	AGC	CCG	CGG	CTA	GCA	CAG	GAT	AGC	CAT	TGT	TGT	GCG	GTA
leu ala g	lv thr	pro	thr	cys	ser	pro	arg	leu	ala	gln	asp	ser	his	cys	cys	ala	val
61/21								91/3	31								
CCC CCA A	AA CGA	TCA	GCC	CTT	CGC	GGA	CAT	GTC	AGC	ACC	CGC	CTT	GGC	CGG	GAG	AGC	GGC
ala pro l	vs arq	ser	ala	leu	arg	gly	his	val	ser	thr	arg	leu	gly	arg	glu	ser	дŢΆ
121/41								151/	51								
GTC GTG A	CC GTG	CTG	TCA	CCA	CGT	CTG	GTT	AGG	CTC	GGG	GCG	CGG	GCT	GGC	GCG	GAG	GAG
val val t	hr val	leu	ser	pro	arg	leu	val	arg	leu	gly	ala	arg	ala	gly	ala	glu	glu
181/61								211.	71								
GTG TGT T	GC GGA	GGA	GGT	GTG	TTG	TAG	TGG	GGA	CGG	CGG	ATC	GGC	CGT	TGG	ACG	CCT	CGG
val cys c	ys gly	gly	gly	val	leu	AMB	trp	gly	arg	arg	ile	gly	arg	trp	thr	pro	arg
241/81								271.	/91								
CCT TGC G	GG ACT	GGG	CAC	ACG	CCG	TCG	TCA	GCG	ATC								
pro cys g	gly thr	gly	his	thr	pro	ser	ser	ala	ile								

SEQ ID No.41B

FIGURE 41B



31/11 TGG CCG GGA CGC CTA CGT GTA GCC CGC GGC TAG CAC AGG ATA GCC ATT GTT GTG CGG TAG trp pro gly arg leu arg val ala arg gly AMB his arg ile ala ile val val arg AMB 91/31 61/21 CGC CAA AAC GAT CAG CCC TTC GCG GAC ATG TCA GCA CCC GCC TTG GCC GGG AGA GCG GCG arg gln asn asp gln pro phe ala asp met ser ala pro ala leu ala gly arg ala ala 151/51 TCG TGA CCG TGC TGT CAC CAC GTC TGG TTA GGC TCG GGG CGC GGG CTG GCG CGG AGG AGG ser OPA pro cys cys his his val trp leu gly ser gly arg gly leu ala arg arg 211/71 TGT GTT GCG GAG GAG GTG TGT TGT AGT GGG GAC GGC GGA TCG GCC GTT GGA CGC CTC GGC cys val ala glu glu val cys cys ser gly asp gly gly ser ala val gly arg leu gly 271/91 241/81 CTT GCG GGA CTG GGC ACA CGC CGT CGT CAG CGA TC leu ala gly leu gly thr arg arg arg gln arg

SEO ID No.41C

FIGURE 41C

Coding sequence Rv2975c predicted by Cole et al, 1998 (Nature 393: 537-544) and containing seq41A

31/11 1/1 gtg ggg acg gcg gat cgg ccg ttg gac gcc tcg gcc ttg cgg gac tgg gca cac gcc gtc val gly thr ala asp arg pro leu asp ala ser ala leu arg asp trp ala his ala val 91/31 gtc agc gat ctg atc ctc cac atc gac gag atc aac cgg ctc aat gtg ttc ccg gtc gct val ser asp leu ile leu his ile asp glu ile asn arg leu asn val phe pro val ala 151/51 gac too gat acc ggc gtc aac atg ctg ttc acc atg cgt gcc gcg gtc gta gaa gct gat asp ser asp thr gly val asn met leu phe thr met arg ala ala val val glu ala asp 211/71 ttg cac gcg aat tcg cag gct gac gcc gaa gac gtg gcg cgg gtt gcg gcc gct ctc gcg leu his ala asn ser gln ala asp ala glu asp val ala arg val ala ala ala leu ala 241/81 gcc ggc gcg cgt tga ala gly ala arg OPA

SEO ID No.41D

FIGURE 41D



ORF according to Cole et al, 1998 (Nature 393: 537-544) and containing Rv2975c

31/111/1 tag get egg gge geg gge tgg ege gga gga ggt gtg ttg egg agg agg tgt gtt gta gtg AMB ala arg gly ala gly trp arg gly gly gly val leu arg arg arg cys val val 91/31 61/21 ggg acg gcg gat cgg ccg ttg gac gcc tcg gcc ttg cgg gac tgg gca cac gcc gtc gtc gly thr ala asp arg pro leu asp ala ser ala leu arg asp trp ala his ala val val 151/51 121/41 age gat ctg atc ctc cac atc gac gag atc aac cgg ctc aat gtg ttc ccg gtc gct gac ser asp leu ile leu his ile asp glu ile asn arg leu asn val phe pro val ala asp 211/71 181/61 tcc gat acc ggc gtc aac atg ctg ttc acc atg cgt gcc gcg gtc gta gaa gct gat ttg ser asp thr gly val asn met leu phe thr met arg ala ala val val glu ala asp leu 271/91 cac gcg aat tcg cag gct gac gcc gaa gac gtg gcg cgg gtt gcg gcc gct ctc gcg gcc his ala asn ser gln ala asp ala glu asp val ala arg val ala ala ala leu ala ala 301/101 ggc gcg cgt tga gly ala arg OPA

SEO ID No.41F

FIGURE 41F

sequence Rv 2974C predicted by Cole et al. (Nature 393:537-544) and which may be in the same reading frame as Seq41D. The sequencing of this region reveals, in one case out of three, a deletion of two nucleotides putting in phase observed in

31/11 1/1 ttg aac gga gct cgc ggc aac tcc ggc gtg atc ctg tcc cag atc ctg cgc ggg atc gca leu asn gly ala arg gly asn ser gly val ile leu ser gln ile leu arg gly ile ala 91/31 gag gtg acc gcg act gcg gcc gcc tct ggc gcg gta ttg cgg gcg gtc gac gcc aac glu val thr ala thr ala ala ala ala ser gly ala val leu arg ala val asp ala asn 151/51 121/41 ged etc ggg ged geg ttg tgg ege gge gtc gag ttg gtc gtc geg teg atg ggt ggc gtg ala leu gly ala ala leu trp arg gly val glu leu val val ala ser met gly gly val 211/71 181/61 gag gtg ccg gga act atc gtc tcg gtg ctg cgg gcc gcc gcc gga gcc gtc gac cag tgc glu val pro gly thr ile val ser val leu arg ala ala gly ala val asp gln cys 271/91 241/81 gcg cac gag ggg ttg gcc ggt gcg gtc acc gcc gcc ggt gac gcg gcg gtc atc gcg ctg ala his glu gly leu ala gly ala val thr ala ala gly asp ala ala val ile ala leu 331/111 301/101 gaa aag acc ccc gaa cag ctt gac gtg ctc gcc gat gcg ggc gcg gtg gac gcc ggc gga glu lys thr pro glu gln leu asp val leu ala asp ala gly ala val asp ala gly gly

SEO ID No.41S

FIGURE 41S

133/185

cog sign can be set gets gets of the case got tag of the case and tage got	361/121				391/131						
arg gly leu leu val leu leu asp ala leu arg ser thr lie cys gly gln ala pro ala 451/151 cgg geg gt to tac gaa coc tog coc geg tog coc geg geg ged acc gac acc go coc arg ala val tyr glu pro ser pro arg ala leu pro thr asp thr ala thr gln arg pro 511/171 gcc ccg caa ttc gag gtg at tat ctg ttg geg ged gta tgd gca geg geg gac cag ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala ala asp gln 541/181 ttg cgg gat cga ctc aag gaa ttg ggt gag tcg gtg gcc atc gcc gct gct ccg acc gac leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 631/211 agc tac tcc gta cac gtc cac acc gac gac gcc ggt gcc gcc gtg gac ggt gre gr yrs ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/221 gtg ggg cg gt tagc cgg atc gta gt gta ctc gcg gt gcc gtg gac gg gt tg ggg gcg gt gtg gcg gt gtg gac acc acc gcc gct gct gcg gcg ggg gcc gc gtg gcg gc	caa aac cta cta	gtt ctg	ctg gac	gcg ttg	g cgc tcc	acc at	c tgc	ggg c	ag gca	cct	gcc
421/141 cgg ggg gtc tac gaa ccc tcg ccg ccg ccg ccg ttg ccg acc gac acc gac acc car ala val try glu pro ser pro arg ala leu pro thr asp thr ala thr gln arg pro 511/171 gcc ccg caa ttc gag gtg atg tat ctg ttg gcg gta tgt gac gca gcg gcg acc ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala asp gln 541/181 ttg ccg gat cga ctc aag gaa ttg ggt gag tcg gtg gcg gtc gcc gcc gac leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala ala pro pro asp 601/201 agc tac tcc gta cac gtc cac acc gac gac gcc gcg gtg gcc gcc gtg gaa gcc gga ttg gcg ser tyr ser val his val his thr asp asp ala ala val ala val glu ala gly leu ala 661/221 gtg ggg ccg gtt acc gcg act gcg gtc gcc gcc gtg gaa gcc gga ttg gcg ser tyr ser val his val his thr asp asp ala cly ala val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 721/241 gcc ggt gcc tcg acc gcg gcc gcc gcc gcc gtc gtc gcc gcc g	arg gly leu leu	val leu	leu asp	ala le	arg ser	thr il	Le cys	gly g	ln ala	pro	ala
arg ala val tyr glu pro ser pro arg ala leu pro thr asp thr ala thr gln arg pro 511/171 gcc ccg caa ttc gag gtg atg tat ctg ttg gcg gta tgt gat gcg gac cag ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala asp gln 541/181 ttg ccg gat ccg act caa gag at ttg gtg gag tcg gcc atc gcc gcc gac leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 601/201 agc tac tcc gta cae gtc caa acc gac gac gcg ggt gcc gcc gtg gaa gcc gga ttg gcg gsr tyr ser val his val his thr asp asp ala glu ala glu ala gly leu ala 661/221 gtg ggc ga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc gat gtg gcg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gcc ggt ggt ga acc gcg ggc gcc gcc gcc gcg gtg gtc gcc gc	421/141				451/151						
goc cog caa ttc gag gtg atg tat cts tts gog gta tgt gat gca gcg gcc gac ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala ala asp gln 541/101 ttg cgg gat cga ctc aag gaa ttg ggt gg gtg ggc atc gcg gct gcc gcc gac leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 651/201 agc tac tcc gta cac gtc cac acc gac gac gcc ggt gcc gcc gtg gag gcc ggt gr gr y ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/221 gtg ggg cga gtt agc cgg atc gtg atc tcg gcg gtg gcc gcc gtg gag gcc ggt gtg ggg gg t agc ggt gtg ggg gg gg gcc gtg gtg ggg gg gg gg gg gg gg gg gg gg	cgg gcg gtc tac	gaa ccc	tcg ccg	cdc dcd	g ttg ccg	acc ga	ac acg	gct a	cc caa	cgc	CCC
gcc ccg caa ttc gag gtg atg tat ctg ttg gcg gta tgt gtg atg ctg cag gac gag gac aag ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala ala asp gln 541/181 ttg cgg gta cga ctc aag gaa ttg ggt gag tcg gtg gcc atc gcc gct ccg ccc gac leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 631/201 agc tac tcc gta cac gtc cac acc gac gac gcg ggt gcc ggt gca gcc ggt ggc gar ty gcg ser tyr ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/221 gtg ggg cga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc gat gtg ggg ga gt agc ggg gcc ggc gtg gcc ggt tcg gga acc agc gat gtg gcg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gcc ggt ggc tgg acg ggg ggc gcc ctg gcc gtg ctg gtc gtc g	arg ala val tyr	glu pro	ser pro	arg ala	a leu pro	thr as	sp thr	ala t	nr gin	arg	pro
sla pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala ala ala ala ala ala ala sal sal	481/161					+~+ ~-	-t aat	aca a	ca aca	asc.	cac
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ttg ogg gat ogg ctc aag gaa ttg ggt gag tog gtg gcc atc gcc gct gct coc ccc ccc ccc leu arg aap arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 601/201 agc tac toc gta cac gtc cac acc gac gac gcc ggt gcc gcc gtg gac gcc ggt gtg gcc gcc gtg gac gcc ggt ser tyr ser val his val his thr asp asp ala qly ala ala val glu ala gly leu ala 661/221 gtg ggg cga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc ggt tg gcg gcc gtg tcc ggg acc agc ggt tg gcg gcc gtg leu gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gcc ggt ggc tgg acg cgg gcc gc gcc gcc gtg tcc gcg gtg gac agc ggt gcc gcc gala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp gly ala ala 811/271 gag ctg ttc gcc ggg gac ggc gcc gcc gtg tcg cgc gcc ggt cac gac ggt gcc gcc glu leu phe ala gly glu gly ala cys val leu arg pro gly pro asp ala val thr pro 841/281 gcc gcc gat atc agt gcc cac cac gct gtg ggg gcc gtg gtg gac acc ggc gcc gcc gla ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala ala pso)01/301 gtg atg gtg ctg ccc aat ggc gta gtc gcc gcc gaa gac ctg gtg gcc gcc gac gac gtg tta gcc gcc yal acc acc gcc yal ala ala glu glu leu val ala glu		e giu vai	met tyr	Ten Te	571/191	cys as	sp ara	ara a	ia ala	шББ	9
leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala pro pro asp 601/201 agc tac tee gta cac gto cac ace gae gae gee ggt gee gee gtg gaa gee ggat tyg cg ser tyr ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/221 gtg ggg cga gtt agc ggg atc gtg ggg ggg ggg ggg ggat gtg ggg ggg gg	541/181	ctc aad	daa ttd	aat aa		acc at	tc acc	act a	ct ccq	ccc	gac
age tac tee gat cae get cae ace gas gas gee get gas gee get gas gee ggt gaa gee gga ttg gee ger tyr ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/221 gtg ggg cga gtt age cgg ate gtg ate tee ggg ete ggt tee ggg ace age gga ttg ceg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gee ggt gge tgg acg cgg gge cgc gtg gtg gtg gtg gt gtg gac ggc gc ala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp gly ala ala ala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp gly ala ala 811/271 gag ctg ttc gcc ggg gag ggc gcc tgc gtg ctg cgc gcg gtg ctg gcg gc	leg egg gat egg	r leu lvs	glu leu	alv al	ı ser val	ala il	le ála	āla ā	la pro	pro	asp
age tac tee gta cae gte cae ace gae gae gee gee gtg gea gee gtg gaa gee gga ttg gee gee gtg gaa gee gga ttg gee gee gtg gaa gee gga ttg gee gae gtg gga gga gta gee gat gtg atc teg gget eee ggt tee ggg ace age gga ttg ceg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gee ggt gge tgg ace ggg gge ege gee gtg etg geg gte gte gae gge gae gg gge gala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp gly ala ala 781/261 gag etg tte gee ggg gag gge etg gae gge gee gtg gtg etg ega egg gge egg glu leu pro asp ala val thr pro 871/291 gee gee gat atc agt gee cae cae egg gtg etg gtg etg gae gge gee gtg leu gtg etg eac ggg gee gee gtg leu ace gg gee gee gae ala ala asp ile ser ala his gln leu val arg pro gly pro asp ala val thr pro 871/291 gee gee gat atc agt gee cae cae etg gtg eeg gee gee gee gee gee gee ge	601/201				631/211						
ser tyr ser val his val his thr asp asp ala gly ala ala val glu ala gly leu ala 661/231 gtg ggg cga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc gga ttg ccg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 721/241	and tac too gta	cac gtc	cac acc	gac ga	c gcc ggt	gcc g	cc gtg	gaa g	cc gga	ttg	gcg
gtg ggg cga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc gga ttg ccg val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gcc ggt ggc tgg acg cgg ggc cgc gcc gtc ctg ggg gtc gtc g	ser tyr ser val	his val	his thr	asp as	p ala gly	ala al	la val	glu a	ıla gly	leu	ala
val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser gly leu pro 751/251 gcc ggt ggc tgg acc cgg ggc cgc gtc gtc gtc gcc gtc gtc g	661/221										
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gcc ggt ggc tgg acg cgg ggc cgc gtc gtg ctg ggc gtc ggc gg		L ser arg	ile val	ile se	r ala leu	gry se	er gry	CHI S	er gry	leu	pro
ala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp gly ala ala 811/271 gag ctg ttc gcc ggg gag ggc gcc tgc gtg ctg cga ccg ggt cca gac gcc gtg aca ccg glu leu phe ala gly glu gly ala cys val leu arg pro gly pro asp ala val thr pro 841/281 gcc gcc gat atc agt gcc cac cag ctg gtg cgg gcc gtg gta gac acc ggc gcc gca ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg ty val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg gtg gcc ggg ttg acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 991/331 gcc acc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gat gcc gac gtg tg acc ggc gtg ala ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcc gcc ccc gca gac gcc gtc gac gac gcc gtg ala ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1111/371 cgt gcc gcc gct gtg ctc ccc gcg cac gga tcg gtc gcc gca gtg gtc gcc gca gtg gcc gcc gca gtg gtc gcc gcc acc gga tcg gtc gcc gcc acc gga tcg gtc gcc gcc acc gga tcg gtc gcc gcc acc gtg gtc gcc gcc gtg gtc gcc gcc gtg gtc gcc gc	721/241			. ~~~ ~+		ata a	tc dac	ממכ מ	rac dot	acc	acc
gag ctg ttc gcc ggg gag ggc gcc tgc gtg ctg cga ccg ggt cca gac gcc gtg aca ccg glu leu phe ala gly glu gly ala cys val leu arg pro gly pro asp ala val thr pro 841/281 gcc gcc gat atc agt gcc cac cag ctg gtg gg gcc gtg gta gac acc ggc gcc gca ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 991/331 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gac gac ggc ttg acg gcg at leu gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcc gct ggc cgt gac gac ggc gcc gcc gac gac ggc gcc gac aca arg ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct ccc gg cac ggc gcc gcc gac ggc gcc gcc	gcc ggt ggc tgg	g acg cgg	ggc cgc	geege	g ccg gcg 1 leu ala	val v	al asp	ggc g	asp alv	ala	ala
gag ctg ttc gcc ggg gag ggc gcc tgc gtg ctg cgg ccg ggt cca gac gcc gtg aca ccg glu leu phe ala gly glu gly ala cys val leu arg pro gly pro asp ala val thr pro 871/291 gcc gcc gat atc agt gcc cac cag ctg gtg cgg gcc gtg gta gac acc ggc gcc gcg cac ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gac gat cg gtg gcc gat gcc gac gtg dla ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc cag gcc cag gcc gac gac gac		o chr arc	dry ard	ala va	811/271	742		5-1	<u>-</u>		
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gcc gcc gat atc agt gcc cac cag ctg gtg cgc gtg gta gac acc ggc gcc gcc ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg cgc gac gga tcg atg gtc cag ggg ttg ala ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc cgc cag gcc gtc gac gac ggc tac agc atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1111/371 gtg gcc gcc ggt gct tcc cgg cac gga tcg gtg gcc att gcc acc caa aag gcg ctg acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 ttgg gcc ggt acc tgc aag ccg ggc gcc ggc ggc gac ggc ggc g	glu leu phe ala	a gly glu	gly ala	cys va	l leu arg	pro g	ly pro	asp a	ala val	thr	pro
ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gga tcg atg gtg cag ggg ttg lai lie gly trp gly val asp val val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc ccc cag gcc gtc gac gac gcc ata agc atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cat gcc acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1171/381 tgg gcc ggt acc tgc aag ccg ggc gac ggc ttg ggt atc gcg gcc gac gac gac gac gac gac gac ga	841/281				871/291						
ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly ala ala his 901/301 gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gga tcg atg gtg cag ggg ttg lai lie gly trp gly val asp val val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc ccc cag gcc gtc gac gac gcc ata agc atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cat gcc acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1171/381 tgg gcc ggt acc tgc aag ccg ggc gac ggc ttg ggt atc gcg gcc gac gac gac gac gac gac gac ga	gcc gcc gat at	c agt gco	cac cag	g ctg gt	g cgg gcc	gtg g	ta gac	acc c	dac acc	gcg	cac
gtg atg gtg ctg ccc aat ggc tat gtg gcc gca gaa gaa ctg gtg gcc ggg tgt acc gcg val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccc gac gga tcg atg gtg cag ggg ttg ala ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc cgc cag gcc gtc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cac gac ggc atc gac gac gcc tac agc atg gcc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gcc gcc gg gcc gcc ggc gg	ala ala asp il	e ser ala	ı his glr	ı leu va	l arg ala	val v	al asp	thr c	дту ата	aıa	nis
val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly cys thr ala 961/321 gcg atc ggc tgg ggc gtc gac gtg gta cec gtg ccg acc gga tcg atg gtg cag ggg ttg ala ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc ccgc cag gcc gtc gac gac ggc tac agc atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cgc att gcc acc caa aag gcg ctg acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggt ggc ggc ggc ggc ggc ggc gg	901/301						ta ata	acc (aaa tat	acc	aca
gcg atc ggc tgg ggc gtc gac gtg gta ccc gga tcc atc gga tcc atc ggc tgg ggc gtc gac gtg gta ccc gga tcc atc atc ggc tgg ggc gtc gac gtg gta ccc gga tcc atc ggc ggc ttc loss ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc gcc ggt gct tcc cgg cac gga tcc ggc gcc atc acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1171/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggc acc acc aa aag gcc ctg acc arg ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu val leu ile 1231/411 gcc gac gac gtg gtg ct act gcc gcc gcc gcc gcc atc ggc ctg gcc acc gac acc atc gga tcg gtc gac gac gtc gtc gcc gcc gcc atc gcc acc acc acc gga tcg gcc acc gac gac gac gac gac gac gac yacc acc acc acc acc gga tcg gcc acc gcc gcc gcc acc gcc gcc gcc acc gcc g	gtg atg gtg ct	g ccc aat	ggc tat	grg go	c gcc gaa	gaa C	en val	ala d	alv cvs	thr	ala
gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gga tcg atg gtg cag ggg ttg ala ile gly trp gly val asp val val pro val pro thr gly ser met val gln gly leu 1021/341 gcc gcg ctg gcc gtg cat gac gcg gcc cgc cag gcc gtc gac gac ggc tac agc atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtc gcc att gcc acc caa aag gcg ctg arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggc gac ggc ggc atc ggc gac ggc gac ggc gac ggc gac ggc gac ggc atc ggc ggc gac ggc ggc atc ggc ggc atc ggc ggc gac ggc atc ggc ggc gac ggc ggc gac ggc ggc atc ggc ggc gac ggc gac ggc ggc gac ggc gg		u pro asi	ı gıy cyı	. vai ai	991/331	giu i	.cu var	u_u ;	5-1 -1-		
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gcc gcg ctg gcc gtg cat gac gcg gcc ccg cag gcc gtc cag gcc gtc gac gac gcc atc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cct gcc gcc gct gct tcc ccg cac gca tcc gtc gcc atc gcc acc caa aag gcc ctg acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tct gcc gct gct tcc cc gcc gcc gcc gcc g	ala ile gly tr	n glv val	L asp val	l val pr	o val pro	thr g	ly ser	met v	val gln	gly	leu
gcc gcg ctg gcc gtg cat gac gcg gcc cgc cag gcc gtc gac gac ggc tac age atg gcc ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr ser met ala 1081/361 cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cgc att gcc acc caa aag gcg ctg acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg gt etg gg ggc gac gag gtg ctg atc trp ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu val leu ile 1201/401 gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gac gcg ctg ctg atc ggg gtc gag tcg	1021/341				1051/35	1					
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cgt gcc gcc ggt gct tcc cgg cac gga tcg gtg cgc att gcc acc caa aag gcg ctg acc arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala leu thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggt atc gcg ggc gac gag gtg ctg atc trp ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu val leu ile 1201/401 gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 gc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gac gtg gct gtc gtc gtc gtc gtc gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc ggc ctg atc ggg gtc gac ggc ctg acc gac cac cat cga ggc gtc gac cac cac cac ggc ggc gta acc gac cac cac cac ggc ggc gac gac ga	ala ala leu al	a val hi:	s asp ala	a ala ar	g gln ala	val a	sp asp	gly 1	tyr ser	met	ala
arg ala ala gly ala ser arg his gly ser val arg lie ala thr glh lys ald led thr 1141/381 tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggt atc gcg ggc gac gag gtg ctg atc trp ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu val leu ile 1201/401 gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac ggc ctg ctg atc ggg gtc gag tag	1081/361				1111/37	1			224 440	. cta	3.00
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tgg gcc ggt acc tgc aag ccg ggc gac ggt ctg ggt atc gcg ggc gac gag gtg ctg atc trp ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu val leu ile 1201/401 gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc gtc gtg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc ggc gtg ctg ctg ctg atc ggg gtc gag tag		y ala se	r arg hi	s gry se	er val arg) 11e a	ila CIII	giii .	rys are	. 104	CIII
trp ala gly thr cys lys pro gly asp gly leu gly leu gly asp glu var leu leu 1201/401 gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg ctg atc ggg gtc gag tag	1141/381	~ +~~ ~~	~ ~~~ ~~	ם מפכ מנ	t cta aat	atc d	ica aac	σac (gag gto	ctq	atc
gc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg ctg atc ggg gtc gag tag	tgg gcc ggt ac	r cue lu	g ccg gg	v asp gl	v leu alv	; ile a	ala gly	asp	glu val	leu	ile
gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg gca tcg gga val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg ctg atc ggg gtc gag tag	1201/401				1721/41	i, L					
val ala asp asp val ala ala ala ala ala ile gly leu val asp leu leu leu ala ser gly 1261/421 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	מדם מכם מפר מפ	t atc ac	c gcg gc	g gcc at	c ggt ctg	g gtc g	gac ctg	ttg	ttg gca	ı tcg	gga
1291/431 ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct gtc ctg gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	val ala asp as	p val al	a ala al	a ala il	Le gly lev	ı val a	asp leu	leu	leu ala	ser	gly
gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala val val leu 1321/441 gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	1261/421				1291/43	31					
1321/441 gaa cgg cat gtg cac gac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	ggc gat ctg gt	g acg gt	g cta at	t ggc g	cc ggc gta	acc g	gaa gac	gtg	gct gto	guc	leu
gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc acc gga cac glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly his 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag		l thr va	l leu il	e giy a.	La gly val	L thr 9	gru asp	Val	ala val	L Vai	. ICa
glu arg his val his asp his his pro gly thr glu leu val ser tyr arg thr gly HIS 1381/461 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	1321/441			+ 662 ~			ata taa	tac	cgc acc	: aaa	cac
1381/461 1411/471 cgc ggc gac gcg ctg ctg atc ggg gtc gag tag	gaa cgg cat gt	g cac ga	c cac ca n hie hi	s pro d	ly thrali	ı leu v	val ser	tyr	arg th	r gly	, his
cgc ggc gac gcg ctg ctg atc ggg gtc gag tag		T HTS GE	F 1172 117	s pro 9.	1411/47	71		_	-		
arg gly asp ala leu leu ile gly val glu AMB	cac aac asc ac	a cta ct	q atc qq	g gtc g							
	arg gly asp al	a leu le	u ile gl	y val g	lu AMB						

SEQ ID No.41S (continued)

FIGURE 41S (continued)

REPLACEMENT SHEET (RULE 26)



Seq41T comprising seq 41F and seq 41S

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31/11
1/1
tta ggc tcg ggg cgc ggg ctg gcg cgg agg agg tgt gtt gcg gag gag gtg tgt tgt agt
leu gly ser gly arg gly leu ala arg arg cys val ala glu glu val cys cys ser
AMB ala arg gly ala gly trp arg gly gly val leu arg arg arg cys val val val
  arg leu gly ala arg ala gly ala glu glu val cys cys gly gly gly val leu AMB trp
                                        91/31
61/21
ggg gac ggc gga tcg gcc gtt gga cgc ctc ggc ctt gcg gga ctg ggc aca cgc cgt cgt
gly asp gly gly ser ala val gly arg leu gly leu ala gly leu gly thr arg arg
 gly thr ala asp arg pro leu asp ala ser ala leu arg asp trp ala his ala val val
  gly arg arg ile gly arg trp thr pro arg pro cys gly thr gly his thr pro ser ser
                                         151/51
cag cga tot gat cot coa cat cga cga gat caa ccg got caa tgt gtt coc ggt cgc tga
gln arg ser asp pro pro his arg arg asp gln pro ala gln cys val pro gly arg OPA
 ser asp leu ile leu his ile asp glu ile asp arg leu asp val phe pro val ala asp
  ala ile OPA ser ser thr ser thr arg ser thr gly ser met cys ser arg ser leu thr
                                         211/71
181/61
ctc cga tac cgg cgt caa cat gct gtt cac cat gcg tgc cgc ggt cgt aga agc tga ttt
leu arg tyr arg arg gln his ala val his his ala cys arg gly arg arg ser OPA phe
 ser asp thr gly val asn met leu phe thr met arg ala ala val val glu ala asp leu
  pro ile pro ala ser thr cys cys ser pro cys val pro arg ser AMB lys leu ile cys
                                         271/91
gca cgc gaa ttc gca ggc tga cgc cga aga cgt ggc gcg ggt tgc ggc cgc tct cgc ggc
ala arg glu phe ala gly OPA arg arg arg gly ala gly cys gly arg ser arg gly
his ala asn ser gln ala asp ala glu asp val ala arg val ala ala ala leu ala ala
  thr arg ile arg arg leu thr pro lys thr trp arg gly leu arg pro leu ser arg pro
                                         331/111
301/101
cgg cgc gcg ttg aac gga gct cgc ggc aac tcc ggc gtg atc ctg tcc cag atc ctg cgc
arg arg ala leu asn gly ala arg gly asn ser gly val ile leu ser gln ile leu arg
 gly ala arg OPA thr glu leu ala ala thr pro ala OPA ser cys pro arg ser cys ala
  ala arg val glu arg ser ser arg gln leu arg arg asp pro val pro asp pro ala arg
                                         391/131
361/121
ggg atc gca gag gtg acc gcg act gcg gcc gcc tct ggc gcg gta ttg cgg gcg gtc
gly ile ala glu val thr ala thr ala ala ala ala ser gly ala val leu arg ala val
 gly ser gln arg OPA pro arg leu arg pro pro leu ala arg tyr cys gly arg ser
  asp arg arg gly asp arg asp cys gly arg arg leu trp arg gly ile ala gly gly arg
421/141
gac gcc aac gcc ctc ggg gcc gcg ttg tgg cgc ggc gtc gag ttg gtc gtc gcg tcg atg
asp ala asn ala leu gly ala ala leu trp arg gly val glu leu val val ala ser met
 thr pro thr pro ser gly pro arg cys gly ala ala ser ser trp ser ser arg arg trp
  arg gln arg pro arg gly arg val val ala arg arg arg val gly arg arg val asp gly
                                         511/171
ggt ggc gtg gag gtg ccg gga act atc gtc tcg gtg ctg cgg gcc gcc gcc gga gcc gtc
gly gly val glu val pro gly thr ile val ser val leu arg ala ala gly ala val
 val ala trp arg cys arg glu leu ser ser arg cys cys gly pro pro pro glu pro ser
  trp arg gly gly ala gly asn tyr arg leu gly ala ala gly arg arg ser arg arg
                                         571/191
541/181
gac cag tgc gcg cac gag ggg ttg gcc ggt gcg gtc acc gcc gcc ggt gac gcg gcg gtc
asp gln cys ala his glu gly leu ala gly ala val thr ala ala gly asp ala ala val
 thr ser ala arg thr arg gly trp pro val arg ser pro pro val thr arg arg ser
  pro val arg ala arg gly val gly arg cys gly his arg arg arg OPA arg gly gly his
                                         631/211
601/201
atc gcg ctg gaa aag acc ccc gaa cag ctt gac gtg ctc gcc gat gcg ggc gcg gtg gac ile ala leu glu lys thr pro glu gln leu asp val leu ala asp ala gly ala val asp
 ser arg trp lys arg pro pro asn ser leu thr cys ser pro met arg ala arg trp thr
   arg ala gly lys asp pro arg thr ala OPA arg ala arg arg cys gly arg gly gly arg
                                         691/231
 661/221
 gcc ggc gga cgg ggc ctg ctg gtt ctg ctg gac gcg ttg cgc tcc acc atc tgc ggg cag
 ala gly gly arg gly leu leu val leu leu asp ala leu arg ser thr ile cys gly gln
 pro ala asp gly ala cys trp phe cys trp thr arg cys ala pro pro ser ala gly arg
   arg arg thr gly pro ala gly ser ala gly arg val ala leu his his leu arg ala gly
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SEQ ID No.41T

FIGURE 41T



751/251 721/241 gca cct gcc cgg gcg gtc tac gaa ccc tcg ccg cgc gcg ttg ccg acc gac acg gct acc ala pro ala arg ala val tyr glu pro ser pro arg ala leu pro thr asp thr ala thr his leu pro gly arg ser thr asn pro arg arg ala arg cys arg pro thr arg leu pro thr cys pro gly gly leu arg thr leu ala ala arg val ala asp arg his gly tyr pro 811/271 781/261 caa cgc ccc gcc ccg caa ttc gag gtg atg tat ctg ttg gcg gta tgt gat gct gca gcg gln arg pro ala pro gln phe glu val met tyr leu leu ala val cys asp ala ala ala asn ala pro pro arg asn ser arg OPA cys ile cys trp arg tyr val met leu gln arg thr pro arg pro ala ile arg gly asp val ser val gly gly met OPA cys cys ser gly 871/291 841/281 gcg gac cag ttg cgg gat cga ctc aag gaa ttg ggt gag tcg gtg gcc atc gcc gct gct ala asp gln leu arg asp arg leu lys glu leu gly glu ser val ala ile ala ala ala arg thr ser cys gly ile asp ser arg asn trp val ser arg trp pro ser pro leu leu gly pro val ala gly ser thr gln gly ile gly OPA val gly gly his arg arg cys ser 931/311 901/301 ccg ccc gac agc tac tcc gta cac gtc cac acc gac gac gcc ggt gcc gcc gtg gaa gcc pro pro asp ser tyr ser val his val his thr asp asp ala gly ala ala val glu ala arg pro thr ala thr pro tyr thr ser thr pro thr thr pro val pro pro trp lys pro ala arg gln leu leu arg thr arg pro his arg arg arg arg cys arg arg gly ser arg 991/331 961/321 gga ttg gcg gtg ggg cga gtt agc cgg atc gtg atc tcg gcg ctc ggt tcc ggg acc agc gly leu ala val gly arg val ser arg ile val ile ser ala leu gly ser gly thr ser asp trp arg trp gly glu leu ala gly ser OPA ser arg arg ser val pro gly pro ala ile gly gly gly ala ser AMB pro asp arg asp leu gly ala arg phe arg asp gln arg 1051/351 1021/341 gly leu pro ala gly gly trp thr arg gly arg ala val leu ala val val asp gly asp asp cys arg pro val ala gly arg gly ala ala pro cys trp arg ser ser thr ala thr ile ala gly arg trp leu asp ala gly pro arg arg ala gly gly arg arg arg arg 1111/371 1081/361 ggt gcc gcc gag ctg ttc gcc ggg gag ggc gcc tgc gtg ctg cga ccg ggt cca gac gcc gly ala ala glu leu phe ala gly glu gly ala cys val leu arg pro gly pro asp ala val pro pro ser cys ser pro gly arg ala pro ala cys cys asp arg val gln thr pro cys arg arg ala val arg arg gly gly arg leu arg ala ala thr gly ser arg arg arg 1171/391 1141/381 gtg aca ccg gcc gcc gat atc agt gcc cac cag ctg gtg cgg gcc gtg gta gac acc ggc val thr pro ala ala asp ile ser ala his gln leu val arg ala val val asp thr gly OPA his arg pro pro ile ser val pro thr ser trp cys gly pro trp AMB thr pro ala asp thr gly arg arg tyr gln cys pro pro ala gly ala gly arg gly arg his arg arg 1231/411 1201/401 gcc gcg cac gtg atg gtg ctg ccc aat ggc tat gtg gcc gcc gaa gaa ctg gtg gcc ggg ala ala his val met val leu pro asn gly tyr val ala ala glu glu leu val ala gly pro arg thr OPA trp cys cys pro met ala met trp pro pro lys asn trp trp pro gly arg ala arg asp gly ala ala gln trp leu cys gly arg arg thr gly gly arg val 1291/431 1261/421 tgt acc gcg gcg atc ggc tgg ggc gtc gac gtg gta ccc gtg ccg acc gga tcg atg gtg cys thr ala ala ile gly trp gly val asp val val pro val pro thr gly ser met val val pro arg arg ser ala gly ala ser thr trp tyr pro cys arg pro asp arg trp cys tyr arg gly asp arg leu gly arg arg gly thr arg ala asp arg ile asp gly ala 1351/451 1321/441 cag ggg ttg gcc gcg ctg gcc gtg cat gac gcg gcc cgc cag gcc gtc gac gac ggc tac gln gly leu ala ala leu ala val his asp ala ala arg gln ala val asp asp gly tyr arg gly trp pro arg trp pro cys met thr arg pro ala arg pro ser thr thr ala thr gly val gly arg ala gly arg ala OPA arg gly pro pro gly arg arg arg leu gln 1411/471 1381/461 age atg gee egt gee ggt get tee egg cae gga teg gtg ege att gee ace caa aag ser met ala arg ala ala gly ala ser arg his gly ser val arg ile ala thr gln lys ala trp pro val pro pro val leu pro gly thr asp arg cys ala leu pro pro lys arg his gly pro cys arg arg cys phe pro ala arg ile gly ala his cys his pro lys gly

SEQ ID No.41T (continued 1)

FIGURE 41T (continued 1)

136/185

1471/491 1441/481 geg etg ace tgg gee ggt ace tge aag eeg gge gae ggt etg ggt ate geg gge gae gag ala leu thr trp ala gly thr cys lys pro gly asp gly leu gly ile ala gly asp glu arg OPA pro gly pro val pro ala ser arg ala thr val trp val ser arg ala thr arg ala asp leu gly arg tyr leu gln ala gly arg arg ser gly tyr arg gly arg arg gly 1531/511 1501/501 gtg ctg atc gtc gcc gac gat gtc gcc gcg gcg gcc atc ggt ctg gtc gac ctg ttg ttg val leu ile val ala asp asp val ala ala ala ala ile gly leu val asp leu leu leu cys OPA ser ser pro thr met ser pro arg arg pro ser val trp ser thr cys cys trp ala asp arg arg arg cys arg arg gly gly his arg ser gly arg pro val val gly 1591/531 gca tcg gga ggc gat ctg gtg acg gtg cta att ggc gcc ggc gta acc gaa gac gtg gct ala ser gly gly asp leu val thr val leu ile gly ala gly val thr glu asp val ala his arg glu ala ile trp OPA arg cys OCH leu ala pro ala OCH pro lys thr trp leu ile gly arg arg ser gly asp gly ala asn trp arg arg arg arg arg arg gly cys 1651/551 1621/541 gtc gtc ctg gaa cgg cat gtg cac gac cac cat cca ggc acc gag ctg gtc tcc tac cgc val val leu glu arg his val his asp his his pro gly thr glu leu val ser tyr arg ser ser trp asn gly met cys thr thr thr ile gln ala pro ser trp ser pro thr ala arg pro gly thr ala cys ala arg pro pro ser arg his arg ala gly leu leu pro his 1711/571 1681/561 acc gga cac cgc ggc gac gcg ctg ctg atc ggg gtc gag tag thr gly his arg gly asp ala leu leu ile gly val glu AMB pro asp thr ala ala thr arg cys OPA ser gly ser ser arg thr pro arg arg ala ala asp arg gly arg val

SEQ ID No.41T (continued 2)

FIGURE 41T (continued 2)

1/1						31/11								
GCC GGT AAC	CC GCG	TCC CA	AG TGC	TAT	CCG	TCC GCC	GGA	CCG	CCC	GAA	ACA	TCA	GCG	GCG
ala gly asn a	ala ala	ser g	ln cys	tyr	pro	ser ala	gly	pro	pro	glu	thr	ser	ala	ala
61/21						91/31								
GGC GCC CCG G	TC GGC	CGC G	GC CGG	GCT	CGA	CCC GCT	CCA	CCT	GGC	CAT	CAG	CGA	CCA	GGT
gly ala pro v	al glv	arg g	ly arg	ala	arg	pro ala	pro	pro	gly	his	gln	arg	pro	gly
121/41						151/51								
ጥአጥ ሮርኔ ርርጥ (GA AGC	GGA C	GG TGT	TGG	GAT	GCA CGC	CCA	ACT	TGC	CGG	CGA	TCG	CGG	CGA
tyr arg gly	ılv ser	glv a	rq cys	trp	asp	ala arg	pro	thr	cys	arg	arg	ser	arg	arg
181/61						211/71								
ጥርር ጥርል ጥርር (GAA CCC	GCG A	CG CAC	ACA	ATG	CCC GCA	GCA	CCG	CAC	GAC	GGC	GCC	CCA	CCG
cys ser ser	ilu pro	ala t	hr his	thr	met	pro ala	ala	pro	his	asp	gly	ala	pro	pro
2/1/81						271/91								
GCT CTT GCA (GTG ACC	TGA T	GA TGA	CAC	TCA	CCC CCA	TAA	GGC	TCG	TCG	GCT	GCG	CCT	GAG
ala leu ala	val thr	OPA O	PA OPA	his	ser	pro pro	OCH	gly	ser	ser	ala	ala	pro	glu
301/101						331/111								
CAA TGC AGT	AAG TTT	ACA C	AA ACG	GAC	TTG	TAA AAA	CCT	GCG	GAG	GTG	GGG	TCT	ATG	GCC
qln cys ser	lvs phe	thr q	ıln thr	asp	leu	OCH lys	pro	ala	glu	val	gly	ser	met	ala
361/121	-1- F	,	*	-		391/131								
AAC AAA CGT	GGC AAT	GCC G	GG CAG	CCT	CTG	CCC TTG	TCG	GAT	С					
asn lys arg	σlv asn	ala d	ılv aln	pro	leu	pro leu	ser	asp						
ash rys arg	9-1 00)— <u>1</u> 5	•		-								

SEQ ID No.42A

FIGURE 42A



1 31/11 1/1 CCG GTA ACG CCG CGT CCC AGT GCT ATC CGT CCG CCG GAC CGC CCG AAA CAT CAG CGG CGG pro val thr pro arg pro ser ala ile arg pro pro asp arg pro lys his gln arg arg 91/31 GCG CCC CGG TCG GCC GCG GCC GGG CTC GAC CCG CTC CAC CTG GCC ATC AGC GAC CAG GTT ala pro arg ser ala ala ala gly leu asp pro leu his leu ala ile ser asp gln val 151/51 ATC GAG GTG GAA GCG GAC GGT GTT GGG ATG CAC GCC CAA CTT GCC GGC GAT CGC GGC GAT ile glu val glu ala asp gly val gly met his ala gln leu ala gly asp arg gly asp 211/71 GCT CAT CGG AAC CCG CGA CGC ACA CAA TGC CCG CAG CAC CGC ACG ACG GCG CCC CAC CGG ala his arg asn pro arg arg thr gln cys pro gln his arg thr thr ala pro his arg 271/91 CTC TTG CAG TGA CCT GAT GAC ACT CAC CCC CAT AAG GCT CGT CGG CTG CGC CTG AGC leu leu gln OPA pro asp asp thr his pro his lys ala arg arg leu arg leu ser 331/111 301/101 AAT GCA GTA AGT TTA CAC AAA CGG ACT TGT AAA AAC CTG CGG AGG TGG GGT CTA TGG CCA asn ala val ser leu his lys arg thr cys lys asn leu arg arg trp gly leu trp pro 391/131 ACA AAC GTG GCA ATG CCG GGC AGC CTC TGC CCT TGT CGG ATC thr asn val ala met pro gly ser leu cys pro cys arg ile

SEQ ID No.42B

FIGURE 42B

31/11 1/1 CGG TAA CGC CGC GTC CCA GTG CTA TCC GTC CGC CGG ACC GCC CGA AAC ATC AGC GGC GGG arg OCH arg arg val pro val leu ser val arg arg thr ala arg asn ile ser gly gly 91/31 61/21 CGC CCC GGT CGG CCG CGG CCG GGC TCG ACC CGC TCC ACC TGG CCA TCA GCG ACC AGG TTA arg pro gly arg pro arg pro gly ser thr arg ser thr trp pro ser ala thr arg leu 151/51 121/41 TCG AGG TGG AAG CGG ACG GTG TTG GGA TGC ACG CCC AAC TTG CCG GCG ATC GCG GCG ATG ser arg trp lys arg thr val leu gly cys thr pro asn leu pro ala ile ala ala met 211/71 CTC ATC GGA ACC CGC GAC GCA CAC AAT GCC CGC AGC ACC GCA CGA CGG CGC CCC ACC GGC leu ile gly thr arg asp ala his asn ala arg ser thr ala arg arg arg pro thr gly 271/91 241/81 TCT TGC AGT GAC CTG ATG ACA CTC ACC CCC ATA AGG CTC GTC GGC TGC GCC TGA GCA ser cys ser asp leu met met thr leu thr pro ile arg leu val gly cys ala OPA ala 331/111 301/101 ATG CAG TAA GTT TAC ACA AAC GGA CTT GTA AAA ACC TGC GGA GGT GGG GTC TAT GGC CAA met gln OCH val tyr thr asn gly leu val lys thr cys gly gly val tyr gly gln 391/131 361/121 CAA ACG TGG CAA TGC CGG GCA GCC TCT GCC CTT GTC GGA TC gln thr trp gln cys arg ala ala ser ala leu val gly

SEQ ID No.42C

FIGURE 42C



Coding sequence Rv2622 predicted by Cole et al., 1998 (Nature 393:537-544) and containing seq42A:

```
31/11
1/1
atg gcc aac aaa cgt ggc aat gcc ggg cag cct ctg ccc ttg tcg gat cga gac gac
Met ala asn lys arg gly asn ala gly gln pro leu pro leu ser asp arg asp asp
                                        91/31
cac atg cag ggg cac tgg ctg ctg gcc cgg ctg ggc aag cgg gtg ctg cgt ccc ggc ggc
his met gln gly his trp leu leu ala arg leu gly lys arg val leu arg pro gly gly
                                        151/51
gto gaa oto acc egg aca otg otg gcc egc gcc gag gtg acc gac gcc gac gtg otc gag
val glu leu thr arg thr leu leu ala arg ala glu val thr asp ala asp val leu glu
                                        211/71
181/61
ctg gca ccg ggc ctg ggc cgc acc gca gcc gaa atc ttg gcc cgc aac ccg cgg tcg tac
leu ala pro gly leu gly arg thr ala ala glu ile leu ala arg asn pro arg ser tyr
                                        271/91
gtg ggg geg gag agc gat ccc aac gcg gcc aac ctg gtc cga cac gtt ctc gcc ggc cgc
val gly ala glu ser asp pro asn ala ala asn leu val arg his val leu ala gly arg
                                        331/111
301/101
ggc gac gtc egg gtc acc gac gcg gcc gat acc gga tta tec gac gcc agc gcc gat gtc
gly asp val arg val thr asp ala ala asp thr gly leu ser asp ala ser ala asp val
                                         391/131
361/121
gte ate gge gag geg atg etg ace atg caa gge aae geg get aaa eae aeg ate gte gee
val ile gly glu ala met leu thr met gln gly asn ala ala lys his thr ile val ala
                                         451/151
421/141
gag gcg gcg cgg gtg ctg agg ccg ggt ggc cgc tac gcg att cac gaa cta gcg ctg gtg
glu ala ala arg val leu arg pro gly gly arg tyr ala ile his glu leu ala leu val
                                         511/171
481/161
ccg gac gac gtc gca gag cag gtc cgc acc gac ctg cgg cag tcg ctg gcc cgc gcg ctc
pro asp asp val ala glu gln val arg thr asp leu arg gln ser leu ala arg ala leu
                                         571/191
541/181
aag gtc aat gcg cgt ccg ctg acc gtt gcg gaa tgg tcg cac ctc tta gcg ggc cat gga
lys val asn ala arg pro leu thr val ala glu trp ser his leu leu ala gly his gly
                                         631/211
 601/201
ctg gtc gtc gaa cac gtt gtc acc gct tcc atg gcg ttg tta caa ccg cga cgg gtg atc
 leu val val glu his val val thr ala ser met ala leu leu gln pro arg arg val ile
                                         691/231
 661/221
 get gae gaa gge ete etg ggt geg etg egg tte gee gga aac etg ete ate eat egt gee
 ala asp glu gly leu leu gly ala leu arg phe ala gly asn leu leu ile his arg ala
                                         751/251
 721/241
 gcg cgt cgg cga gtc ctg ttg atg cgc cac aca ttc cgc agg cat cgt gaa cgc ttg aca
 ala arg arg arg val leu leu met arg his thr phe arg arg his arg glu arg leu thr
                                         811/271
 781/261
 gee gte gee att gte geg cae aaa eeg eae gte gat teg tga
 ala val ala ile val ala his lys pro his val asp ser OPA
```

SEQ ID No.42D

FIGURE 42D



ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv2622

```
31/11
1/1
taa aaa cct gcg gag gtg ggg tct atg gcc aac aaa cgt ggc aat gcc ggg cag cct ctg
OCH lys pro ala glu val gly ser met ala asn lys arg gly asn ala gly gln pro leu
                                        91/31
ccc ttg tcg gat cga gac gac cac atg cag ggg cac tgg ctg ctg gcc cgg ctg ggc
pro leu ser asp arg asp asp his met gln gly his trp leu leu ala arg leu gly
                                        151/51
121/41
aag cgg gtg ctg cgt ccc ggc ggc gtc gaa ctc acc cgg aca ctg ctg gcc cgc gag
lys arg val leu arg pro gly gly val glu leu thr arg thr leu leu ala arg ala glu
                                        211/71
gtg acc gac gcc gac gtg ctc gag ctg gca ccg ggc ctg ggc cgc acc gca gcc gaa atc
val thr asp ala asp val leu glu leu ala pro gly leu gly arg thr ala ala glu ile
                                        271/91
241/81
ttq gcc cgc aac ccg cgg tcg tac gtg ggg gcg gag agc gat ccc aac gcg gcc aac ctg
leu ala arg asn pro arg ser tyr val gly ala glu ser asp pro asn ala ala asn leu
                                        331/111
301/101
qtc cga cac gtt ctc gcc ggc cgc ggc gac gtc cgg gtc acc gac gcg gcc gat acc gga
val arg his val leu ala gly arg gly asp val arg val thr asp ala ala asp thr gly
                                        391/131
361/121
tta tcc gac gcc agc gcc gat gtc gtc atc ggc gag gcg atg ctg acc atg caa ggc aac
leu ser asp ala ser ala asp val val ile gly glu ala met leu thr met gln gly asn
                                         451/151
421/141
gcg gct aaa cac acg atc gtc gcc gag gcg gcg gtg ctg agg ccg ggt ggc cgc tac
ala ala lys his thr ile val ala glu ala ala arg val leu arg pro gly gly arg tyr
                                         511/171
481/161
gcg att cac gaa cta gcg ctg gtg ccg gac gac gtc gca gag cag gtc cgc acc gac ctg
ala ile his glu leu ala leu val pro asp asp val ala glu gln val arg thr asp leu
                                         571/191
cgg cag tcg ctg gcc cgc gcg ctc aag gtc aat gcg cgt ccg ctg acc gtt gcg gaa tgg
arg gln ser leu ala arg ala leu lys val asn ala arg pro leu thr val ala glu trp
                                         631/211
 601/201
 teg cae ete tta geg gge cat gga etg gte gte gaa eae gtt gte ace get tee atg geg
 ser his leu leu ala gly his gly leu val val glu his val val thr ala ser met ala
                                         691/231
 ttg tta caa ccg cga cgg gtg atc gct gac gaa ggc ctc ctg ggt gcg ctg cgg ttc gcc
 leu leu gln pro arg arg val ile ala asp glu gly leu leu gly ala leu arg phe ala
                                         751/251
 gga aac ctg ctc atc cat cgt gcc gcg cgt cgg cga gtc ctg ttg atg cgc cac aca ttc
 gly asn leu leu ile his arg ala ala arg arg val leu leu met arg his thr phe
                                         811/271
 cgc agg cat cgt gaa cgc ttg aca gcc gtc gcc att gtc gcg cac aaa ccg cac gtc gat
 arg arg his arg glu arg leu thr ala val ala ile val ala his lys pro his val asp
 841/281
 tcg tga
 ser OPA
```

SEQ ID No.42F

FIGURE 42F



1/1
atc gcg cgt gac atc gat gac cag ggt cgg ctg tgt ctg gac gtc ggc ggt cga acg gta ile ala arg asp ile asp asp gln gly arg leu cys leu asp val gly gly arg thr val 91/31
gtt gtt tca gcg ggc gac gtg gtg cat ttg cgt taa ctc gcg cgg agc tgg cgt ccc caa val val ser ala gly asp val val his leu arg OCH leu ala arg ser trp arg pro gln 121/41
aag att aag gtc gcg ggc atg agc tat ccg gag aat gtc ctg gcc gct ggc gag cag gtc lys ile lys val ala gly met ser tyr pro glu asn val leu ala ala gly glu gln val 181/61
gtt ctg cac cgc cat ccg cac tgg aat cgc tta atc tgg ccc gtc gtg gtg ctg gtc ttg val leu his arg his pro his trp asn arg leu ile trp pro val val val leu val leu 241/81
ctg acc ggg ttg gcg gcg ttc ggg tcc gga ttc gtc acc tgg acc cct tgg cag cag atc leu thr gly leu ala ala phe gly ser gly phe val asn ser thr pro trp gln gln ile

SEQ ID No.43A

FIGURE 43A

31/11 tcg cgc gtg aca tcg atg acc agg gtc ggc tgt gtc tgg acg tcg gcg gtc gaa cgg tag ser arg val thr ser met thr arg val gly cys val trp thr ser ala val glu arg AMB . 91/31 ttg ttt cag cgg gcg acg tgg tgc att tgc gtt aac tcg cgc gga gct ggc gtc ccc aaa leu phe gln arg ala thr trp cys ile cys val asn ser arg gly ala gly val pro lys 151/51 aga tta agg tcg cgg gca tga gct atc cgg aga atg tcc tgg ccg ctg gcg agc agg tcg arg leu arg ser arg ala OPA ala ile arg arg met ser trp pro leu ala ser arg ser 211/71 ttc tgc acc gcc atc cgc act gga atc gct taa tct ggc ccg tcg tgg tgc tgg tct tgc phe cys thr ala ile arg thr gly ile ala OCH ser gly pro ser trp cys trp ser cys 271/91 241/81 tga ccg ggt tgg cgg cgt tcg ggt ccg gat tcg tca act cga cac ctt ggc agc aga tc OPA pro gly trp arg arg ser gly pro asp ser ser thr arg his leu gly ser arg

SEQ ID No.43B

FIGURE 43B

141/185

31/11 1/1 cgc gcg tga cat cga tga cca ggg tcg gct gtg tct gga cgt cgg cgg tcg aac ggt agt arg ala OPA his arg OPA pro gly ser ala val ser gly arg arg ser asn gly ser 91/31 tgt ttc agc ggg cga cgt ggt gca ttt gcg tta act cgc gcg gag ctg gcg tcc cca aaa cys phe ser gly arg arg gly ala phe ala leu thr arg ala glu leu ala ser pro lys 151/51 gat taa ggt cgc ggg cat gag cta tcc gga gaa tgt cct ggc cgc tgg cga gca ggt cgt asp OCH gly arg gly his glu leu ser gly glu cys pro gly arg trp arg ala gly arg 211/71 tet gca ccg cca tcc gca ctg gaa tcg ctt aat ctg gcc cgt cgt ggt gct ggt ctt gct ser ala pro pro ser ala leu glu ser leu asn leu ala arg arg gly ala gly leu ala 271/91 gac cgg gtt ggc ggc gtt cgg gtc cgg att cgt caa ctc gac acc ttg gca gca gat c asp arg val gly gly val arg val arg ile arg gln leu asp thr leu ala ala asp

SEQ ID No.43C

FIGURE 43C

Coding sequence Rv3278c predicted by Cole et al., 1998 (Nature 393:537-544) and containing seq43A:

1/1	31/11
ata ago tat con dag aat oto cto occ oct	ggc gag cag gtc gtt ctg cac cgc cat ccg
Met ser tyr pro glu asn val leu ala ala	gly glu gln val val leu his arg his pro
61 / 21	91/31
and the aat one tha ato the occ etc etc	gtg ctg gtc ttg ctg acc ggg ttg gcg gcg
his tro asn arg leu ile tro pro val val	val leu val leu leu thr gly leu ala ala
101//1	151/51
the aga too aga the ate age tog aca cet	tgg cag cag atc gct aag aac gtg att cac
phe gly ser gly phe val asn ser thr pro	trp gln gln ile ala lys asn val ile his
101/61	211/71
are ato ato tag aga ato tag tta ato ato	gtc ggc tgg ctc acg ctg tgg cca ttc ctg
ala val ile trp glv ile trp leu val ile	e val gly trp leu thr leu trp pro phe leu
041/01	271/91
and the ste acc aca cat the ete ete acc	aac egg egg gtg atg tte egg eat ggt gtg
ser tro leu thr thr his phe val val the	r asn arg arg val met phe arg his gly val
201/101	331/111
ata and and and and ata dad ata dog cta	a gca cgg atc aac agc gtg gag ttc cgg gac
len thr arg ser glv ile asp ile pro len	ala arg ile asn ser val glu phe arg asp
261/121	391/131
and ata the dad end att the end acc que	g acg ttg att atc gag tcc gcg tca caa gat
arg ile phe glu arg ile phe arg thr gly	y thr leu ile ile glu ser ala ser gln asp
421/141	451/151
and ata day the tac age att eeg ege et	g cgg gag gtg cat gcg ttg ctg tat cac gag
pro leu glu phe tvr asn ile pro arg le	u arg glu val his ala leu leu tyr his glu
481/161	511/171
gtt ttc gac acc ctg ggc tcc gac gag tc	g ccc agc tga
val phe asp thr leu gly ser asp glu se	r pro ser OPA

SEQ ID No.43D

FIGURE 43D



ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv3278c

```
31/11
1/1
taa ctc gcg cgg agc tgg cgt ccc caa aag att aag gtc gcg ggc atg agc tat ccg gag
OCH leu ala arg ser trp arg pro gln lys ile lys val ala gly met ser tyr pro glu
                                        91/31
aat gtc ctg gcc gct ggc gag cag gtc gtt ctg cac cgc cat ccg cac tgg aat cgc tta
asn val leu ala ala gly glu gln val val leu his arg his pro his trp asn arg leu
                                        151/51
atc tgg ccc gtc gtg gtg ctg gtc ttg ctg acc ggg ttg gcg gcg ttc ggg tcc gga ttc
ile trp pro val val leu val leu leu thr gly leu ala ala phe gly ser gly phe
                                        211/71
gtc aac tog aca cot tgg cag cag atc gct aag aac gtg att cac gcg gtc atc tgg ggg
val asn ser thr pro trp gln gln ile ala lys asn val ile his ala val ile trp gly
                                        271/91
atc tgg ttg gtg atc gtc ggc tgg ctc acg ctg tgg cca ttc ctg agc tgg ctg acc aca
ile trp leu val ile val gly trp leu thr leu trp pro phe leu ser trp leu thr thr
                                        331/111
301/101
cat ttc gtg gtg acc aac cgg cgg gtg atg ttc cgg cat ggt gtg ctg acc cgc agc ggg
his phe val val thr asn arg arg val met phe arg his gly val leu thr arg ser gly
                                         391/131
361/121
atc gac ata ccg cta gca cgg atc aac agc gtg gag ttc cgg gac cgg atc ttc gag cgg
ile asp ile pro leu ala arg ile asn ser val glu phe arg asp arg ile phe glu arg
                                         451/151
421/141
att ttt cgc acc ggg acg ttg att atc gag tcc gcg tca caa gat ccg ctc gag ttc tac
ile phe arg thr gly thr leu ile ile glu ser ala ser gln asp pro leu glu phe tyr
                                         511/171
481/161
aac att ccg cgc ctg cgg gag gtg cat gcg ttg ctg tat cac gag gtt ttc gac acc ctg
asn ile pro arg leu arg glu val his ala leu leu tyr his glu val phe asp thr leu
541/181
ggc tcc gac gag tcg ccc agc tga
gly ser asp glu ser pro ser OPA
```

SEQ ID No.43F

FIGURE 43F



31/11 1/1 gcc aag atg gat gtc tac caa cgc acc gcc gcc ggc tgg cag ccg ctc aag acc ggt atc ala lys met asp val tyr gln arg thr ala ala gly trp gln pro leu lys thr gly ile 91/31 acc acc cat atc ggt tcg gcg ggc atg gcg ccg gaa gcc aag agc gga tat ccg gcc act thr thr his ile gly ser ala gly met ala pro glu ala lys ser gly tyr pro ala thr 151/51 121/41 ccg atg ggg gtt tac agc ctg gac tcc gct ttt ggc acc gcg ccg aat ccc ggt ggc ggg pro met gly val tyr ser leu asp ser ala phe gly thr ala pro asn pro gly gly 211/71 181/61 ttg ccg tat acc caa gtc gga ccc aat cac tgg tgg agt ggc gac gac aat agc ccc acc leu pro tyr thr gln val gly pro asn his trp trp ser gly asp asp asn ser pro thr 271/91 ttt aac tcc atg cag gtc tgt cag aag tcc cag tgc ccg ttc agc acg gcc gac agc gag phe asn ser met gln val cys gln lys ser gln cys pro phe ser thr ala asp ser glu 331/111 301/101 aac ctg caa atc ccg cag tac aag cat tcg gtc gtg atg ggc gtc aac aag gcc aag gtc asn leu gln ile pro gln tyr lys his ser val val met gly val asn lys ala lys val 391/131 cca ggc aaa ggc tcc gcg ttc ttc ttt cac acc acc gac ggc ggg ccc acc gcg ggt tgt pro gly lys gly ser ala phe phe his thr thr asp gly gly pro thr ala gly cys 421/141 gtg gcg atc val ala ile

SEQ ID No.44A

FIGURE 44A

31/11 1/1 cca aga tgg atg tct acc aac gca ccg ccg ccg gct ggc agc cgc tca aga ccg gta tca pro arg trp met ser thr asn ala pro pro pro ala gly ser arg ser arg pro val ser 91/31 cca ccc ata teg gtt egg egg gca tgg ege egg aag eca aga geg gat ate egg eea ete pro pro ile ser val arg arg ala trp arg arg lys pro arg ala asp ile arg pro leu 151/51 121/41 cga tgg ggg ttt aca gcc tgg act ccg ctt ttg gca ccg cgc cga atc ccg gtg gcg ggt arg trp gly phe thr ala trp thr pro leu leu ala pro arg arg ile pro val ala gly 211/71 tgc cgt ata ccc aag tcg gac cca atc act ggt gga gtg gcg acg aca ata gcc cca cct cys arg ile pro lys ser asp pro ile thr gly gly val ala thr thr ile ala pro pro 271/91 241/81 tta act cca tgc agg tct gtc aga agt ccc agt gcc cgt tca gca cgg ccg aca gcg aga leu thr pro cys arg ser val arg ser pro ser ala arg ser ala arg pro thr ala arg 331/111 acc tgc aaa tcc cgc agt aca agc att cgg tcg tga tgg gcg tca aca agg cca agg tcc thr cys lys ser arg ser thr ser ile arg ser OPA trp ala ser thr arg pro arg ser 391/131 361/121 cag gca aag gct ccg cgt tct tct ttc aca cca ccg acg gcg ggc cca ccg cgg gtt gtg gln ala lys ala pro arg ser ser phe thr pro pro thr ala gly pro pro arg val val 421/141 tgg cga tc trp arg

SEQ ID No.44B

FIGURE 44B

THE A CHERTSHIP CHEEDING CHILL IS AC



31/11 1/1 caa gat gga tgt cta cca acg cac cgc cgc cgg ctg gca gcc gct caa gac cgg tat cac qln asp gly cys leu pro thr his arg arg leu ala ala ala gln asp arg tyr his 61/21 91/31 cac cca tat egg tte ggc ggg cat ggc gcc gga age caa gag egg ata tee gge cae tee his pro tyr arg phe gly gly his gly ala gly ser gln glu arg ile ser gly his ser 151/51 121/41 gat ggg ggt tta cag cct gga ctc cgc ttt tgg cac cgc gcc gaa tcc cgg tgg cgq gtt asp gly gly leu gln pro gly leu arg phe trp his arg ala glu ser arg trp arg val 211/71 181/61 gcc gta tac cca agt cgg acc caa tca ctg gtg gag tgg cga cga caa tag ccc cac ctt ala val tyr pro ser arg thr gln ser leu val glu trp arg arg gln AMB pro his leu 271/91 241/81 taa ctc cat gca ggt ctg tca gaa gtc cca gtg ccc gtt cag cac ggc cga cag cga gaa OCH leu his ala gly leu ser glu val pro val pro val gln his gly arg gln arg glu 331/111 301/101 cct gca aat ccc gca gta caa gca ttc ggt cgt gat ggg cgt caa caa ggc caa ggt ccc pro ala asn pro ala val gln ala phe gly arg asp gly arg gln gln gly gln gly pro 391/131 361/121 agg caa agg ctc ege gtt ctt ctt tca cac cac ega egg egg gee cac ege ggg ttg tgt arg gln arg leu arg val leu leu ser his his arg arg ala his arg gly leu cys 421/141 ggc gat c gly asp

SEQ ID No.44C

FIGURE 44C



Coding sequence Rv0309 predicted by Cole et al., 1998 (Nature 393:537-544) and containing Seq44A:

1/1								31/1	1								
atg agc co	a ctc	cta	act	ttσ	cta	tac	act	aca	gta	tqc	acq	ggc	tgc	gtt	gct	gtg	gtt
Met ser an	ra leu	leu	ala	leu	leu	CVS	ala	ālā	val	cys	thr	gly	cys	val	ala	val	val
61/21								91/3	31								
ctc aca co	ca qtq	aqc	ctg	gcc	gtc	gtc	aac	ccg	tgg	ttc	gcg	aac	tcg	gtc	ggc	aat	gcc
leu ala pi	o val	ser	leu	ala	val	val	asn	pro	trp	phe	ala	asn	ser	val	gly	asn	ala
121/41								151/	51								
act cag gt	g gtt	tcg	gtg	gtg	gga	acc	ggc	ggt	tcg	acg	gcc	aag	atg	gat	gtc	tac	caa
thr gln va	al val	ser	val	val	gly	thr	gly	gly	ser	thr	ala	lys	met	asp	val	tyr	gln
181/61								211/									
cgc acc go	cc gcc	ggc	tgg	cag	ccg	ctc	aag	acc	ggt	atc	acc	acc	cat	atc	ggt	tcg	gcg
arg thr a	la ala	gly	trp	gln	pro	leu	lys	thr	gra	ıle	tnr	tnr	nis	iie	дтĀ	ser	ala
241/81								271,		- at	~~~	2 t a	aaa	a++	+ = C	300	ctc
ggc atg g	cg ccc	gaa	gcc	aag	agc	gga	tat	ccg	gcc	act +hr	nro	mot	999	7727	tur	ear	1 211
gly met a	la pro	giu	ala	Tys	ser	дтХ	cyr	Dro.	/111	CIII	pro	mec	9 ± Y	val	CYL	DCI	
301/101 gac tcc g				~~~	000	22+	ccc			aaa	tta	cca	tat	acc	caa	atc	gga
gac tcc gasp ser a	Ct ttt	ggc	acc	212	nro	aat	nro	al v	al a	alv	leu	pro	tvr	thr	aln	val	alv
_	ra pne	gry	CIII	ala	Pro	asn	PLO	391	/131	9-1		L	- 2		٠.		<i>-</i>
361/121 ccc aat c	a ta	. +	aat	ממכ	aac	gac	aat			acc	ttt	aac	tcc	atq	caq	gtc	tgt
pro asn h	ie tr	tro	ser	alv	asp	asp	asn	ser	pro	thr	phe	asn	ser	met	gln	val	cys
421/141	TO CT	CLP	DCI	9-1	шор	wo P		451	/151		•				-		_
cag aag t	cc cac	r tac	cca	ttc	agc	acq	gcc	qac	agc	gag	aac	ctg	caa	atc	ccg	cag	tac
gln lys s	er alı	cvs	pro	phe	ser	thr	ala	asp	ser	glu	asn	leu	gln	ile	pro	gln	tyr
481/161								511	/171								
aad cat t	ca at	gtg	atg	ggc	gtc	aac	aag	gcc	aag	gtc	cca	ggc	aaa	ggc	tcc	gcg	ttc
lys his s	er va	L val	met	gly	val	asn	lys	ala	lys	val	pro	gly	lys	gly	ser	ala	phe
541/181								571	/191								
ttc ttt c	ac ac	c acc	gac	ggc	ggg	ccc	acc	gcg	ggt	tgt	gtg	gcg	atc	gac	gat	gcc	acg
phe phe h	is th	r thr	asp	gly	gly	pro	thr	ala	gly	cys	val	ala	ıle	asp	asp	ата	Chr
601/201									/211		4_		~+~	. ~~~		+	
ctg gtg c	ag at	c ato	: cgt	tgg	ctg	cgg	cct	ggt	gcg	gtg	atc	geg	ila	, gcc	aay Tue	೧೭೪	•
leu val g	ıln il	e ile	arg	trp	Leu	arg	pro	о атй	ата	val	тте	ard	TTE	. ата	туз	COL	

SEQ ID No.44D

FIGURE 44D



ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv0309

```
31/11
1/1
tga gcg atg agc cga ctc cta gct ttg ctg tgc gct gcg gta tgc acg ggc tgc gtt gct
OPA ala met ser arg leu leu ala leu leu cys ala ala val cys thr gly cys val ala
                                        91/31
gtg gtt ctc gcg cca gtg agc ctg gcc gtc gtc aac ccg tgg ttc gcg aac tcg gtc ggc
val val leu ala pro val ser leu ala val val asn pro trp phe ala asn ser val gly
                                        151/51
aat gcc act cag gtg gtt tcg gtg gtg gga acc ggc ggt tcg acg gcc aag atg gat gtc
asn ala thr gln val val ser val val gly thr gly gly ser thr ala lys met asp val
                                        211/71
181/61
tac caa ege ace gee gee tgg cag eeg ete aag ace ggt ate ace ace cat ate ggt
tyr gln arg thr ala ala gly trp gln pro leu lys thr gly ile thr thr his ile gly
                                        271/91
241/81
teg geg gge atg geg eeg gaa gee aag age gga tat eeg gee aet eeg atg ggg gtt tae
ser ala gly met ala pro glu ala lys ser gly tyr pro ala thr pro met gly val tyr
                                        331/111
301/101
age ctg gac tee get ttt ggc ace geg eeg aat eee ggt gge ggg ttg eeg tat ace eaa
ser leu asp ser ala phe gly thr ala pro asn pro gly gly leu pro tyr thr gln
                                        391/131
361/121
gtc gga ccc aat cac tgg tgg agt ggc gac gac aat agc ccc acc ttt aac tcc atg cag
val gly pro asn his trp trp ser gly asp asp asn ser pro thr phe asn ser met gln
                                         451/151
421/141
gtc tgt cag aag tcc cag tgc ccg ttc agc acg gcc gac agc gag aac ctg caa atc ccg
val cys gln lys ser gln cys pro phe ser thr ala asp ser glu asn leu gln ile pro
                                         511/171
cag tac aag cat tcg gtc gtg atg ggc gtc aac aag gcc aag gtc cca ggc aaa ggc tcc
gln tyr lys his ser val val met gly val asn lys ala lys val pro gly lys gly ser
                                         571/191
541/181
geg tto tto ttt cac acc gac ggc ggg ccc acc gcg ggt tgt gtg gcg atc gac gat
ala phe phe phe his thr thr asp gly gly pro thr ala gly cys val ala ile asp asp
                                         631/211
qcc acg ctg gtg cag atc atc cgt tgg ctg cgg cct ggt gcg gtg atc gcg atc gcc aag
ala thr leu val gln ile ile arg trp leu arg pro gly ala val ile ala ile ala lys
 taa
 OCH
```

SEQ ID No.44F

FIGURE 44F



Cloned fragment fused with phoA

1/1										31/1									
ant.	ata	ccc	gga	cac	caq	gtc	atc	cgg	cga (gat 4	ggt	gat	cga	ggc	tcg	gac	ccg	cag	gca
asp .	leu	pro	gly	his	gln	val	ile	arg	arg .	asp	gly	asp	arg	gly	ser	asp	pro	gln	ala
61/2	1									91/3	1								
tcc	ggt	agc	cag	agg	cac	cag	cat	cag	caa	cat	cgc	gat	ggc	cag	cat	gcc	gcg	ccg	tcg
ser	gly	ser	gln	arg	his	gln	his	gln	gln :	his	arg	asp	дТĀ	gın	his	a⊥a	ата	pro	ser
121/	41									151/	51								
ggt	cct	tgc	cac	tcg	cga	tcc	ttg	gga	tga	cgg	tgg	ggc	ila	gct	agc	90g	hie	aln	val
gly	pro	cys	his	ser	arg	ser	leu	g⊥y	OPA	arg 211/	trp 71	дтУ	тте	ala	ser	ата	1112	gin	vai
181/	61					+~~	000	ata				tca	aac	gcg	aat	tag	aσc	aat	agc
atc	gtg	cca	gac	cgg	gca	cyc	ara	wal	alv	lvs	leu	ser	alv	ala	alv	AMB	ser	gly	ser
		pro	asp	arg	ата	Cys	ary	Val	9-1	271/	91		9-1		<i>J</i> _ 4				
241/	01	~~~	244	ata	aca	aat	act	caa	aaa			aca	aaq	tgg	tag	ccg	cgg	atg	atg
geg	ara	nro	ayy	met	ala	asn	ala	arq	gly	ser	pro	ala	lys	trp	AMB	pro	arg	met	met
301/	101									331/	111								
+ ~ ~	ata	aaq	ccc	aac	cgg	cgg	tac	aac	cgc	cac	gcc	cga	ttg	tcc	tca	ccg	ttg	gtc	tcc
ser	val	lys	pro	asn	arg	arg	tyr	asn	arg	his	ala	arg	leu	ser	ser	pro	leu	val	ser
261/	121									391/	131								
ggt	gtg	gag	agc	agg	acg	ttg	tcc	tcg	tcg	cga	ccg	gct	agc	agt	cgg	cgg	gcc	aac	gcc
gly	val	glu	ser	arg	thr	leu	ser	ser	ser	arg	pro	ala	ser	ser	arg	arg	ala	asii	ara
421/	141									451/		+~~	22+	tas	~+ <i>~</i>	220	tca	aan	tad
tcc	ccg	agg	cca	cgg	cct	tga	gcg	cgg	gga	agg	acg	cgc	aat	tca	1721	aac	ser	lvs	AMB
		arg	pro	arg	pro	OPA	ата	arg	дту	511/	1171	Cys	asii	ser	Val	abii	501	-1-	
481/	161				~~~	ata	act	200	cac			cca	cta	cgt	tac	aaq	ccc	agt	acc
ctg	gtc	atc	agt	cgg	212	ila	ala	ayy	ara	alv	lvs	pro	leu	arg	cvs	lys	pro	ser	thr
	vai /181	116	ser	ary	ата	110	ana	arg	429	571	191	L – -		_	-	-	-		
	+~~	tat	tac	cac	cac	taa	cca	aac	qcc	ccg	gga	tag	ccg	tac	gcc	act	ccg	agc	att
thr	CVS	CVS	cvs	his	his	trp	pro	gly	āla	pro	gly	AMB	pro	tyr	ala	thr	pro	ser	ile
601	/201									631,	/ 211								
		ttg	ctc	agt	tcg	gcg	gcc	gac	ggc	agc	gcc	gtg	gtg	tcg	gcg	gcc	tcg	gcc	tgt
qly	ala	leu	leu	ser	ser	ala	ala	asp	gly	ser	ala	vai	val	ser	ala	ala	ser	ala	cys
661	/221									691	/ 231								
tcg	gct	gcc	gtt	acc	tcg	acg	gcc	gcg	acc	gcc	tgc	cag	ccg	cgc	cgc	cgg	atg	Lgc	tcc
ser	ala	ala	val	thr	ser	thr	ala	ala	thr	ala	Cys	gın	pro	arg	arg	ary	mec	Cys	ser
721	/241								~+~		/251		r tao	י כמכ	ato	aco	r t.ca	aca	tac
agc	cac	att	ggg	gcg	cgc	aaa	gtc	tcg	grg	nro	len	ggg alt	, AME	, cyc Raro	ile	ala	ser	thr	tac tyr
			g gr	а а а а	arg	тys	Val	ser	vai	811	/271	9-1	, , , , , ,	, 429					-
781	/261			+ 4 7 7		. 200	cao	י כמכ	tica				a aac	gac	aga	tcc	ato	ago	g aat
acc	gtc	agg	g gca		nro	, ayy	aro	aro	ser	ile	ser	lei	gly	, gly	arg	sei	met	arg	asn
0.41	/201									0/1	1271	_							
- + -	~~~		e acc	ı cac	ı tat	. cct	cct	cat	gtg	atq	aac	cga	a tgo	gtg	ctt	gcg	g cac	caç	tat tvr
ile	ala	ası	n ala	arc	CVS	pro	pro	his	vaĺ	met	asr	arç	g cys	s val	. leı	ıala	a his	glı	ı tyr
0.01	/301									931	/311	L							
			c cga	a tga	a ggo	cgc	ccg	g cgc	: tgg	acg	gg	g cti	t gta	a gcc	, tat	aão	c cgt	: tt(c cgc
arq	thi	se:	r ar	OP?	A gly	arg	g pro	arç	trp	thr	gly	y le	u val	ı ala	t ty	c dr	y arc	g pne	e arg
_																			

SEQ ID No.45ZA

FIGURE 45ZA

148/185

991/331 961/321 tca gct cgt cgc tgc gcc gcc gcg ata gaa tcg ccc gcg aac cag tgg tac ggc gca ser ala arg arg cys gly ala ala gly ile glu ser pro ala asn gln trp tyr gly ala 1051/351 gat tga cct cgt atc atc tga gtt agt tgc ccg cgc aat ggg cat ccg cgt gtt atc ggt asp OPA pro arg ile ile OPA val ser cys pro arg asn gly his pro arg val ile gly 1111/371 att acg tga cag tct gtc ggc aag gag gga cgc atg cca ctc tcc gat cat gag cag cgg ile thr OPA gln ser val gly lys glu gly arg met pro leu ser asp his glu gln arg 1171/391 atg ctt gac cag atc gag agc gct ctc tac gcc gaa gat ccc aag ttc gca tcg agt gtc met leu asp gln ile glu ser ala leu tyr ala glu asp pro lys phe ala ser ser val 1231/411 cgt ggc ggg ggc ttc cgc gca ccg acc gcg cgg cgc ctg cag ggc gcg gcg ttg ttc arg gly gly phe arg ala pro thr ala arg arg leu gln gly ala ala leu phe 1291/431 atc atc ggt ctg ggg atg ttg gtt tcc ggc gtg gcg ttc aaa gag acc atg atc gga agt ile ile gly leu gly met leu val ser gly val ala phe lys glu thr met ile gly ser 1351/451 1321/441 ttc ccg ata ctc agc gtt ttc ggt ttt gtc gtg atg ttc ggt gtg gtg tat gcc atc phe pro ile leu ser val phe gly phe val val met phe gly gly val val tyr ala ile 1411/471 1381/461 ace ggt cet egg ttg tee ggc agg atg gat egt ggc gga teg get get ggg get teg ege thr gly pro arg leu ser gly arg met asp arg gly gly ser ala ala gly ala ser arg 1471/491 1441/481 cag cgt cgt acc aag ggg gcc ggg ggc tca ttc acc agc cgt atg gaa gat c gln arg arg thr lys gly ala gly gly ser phe thr ser arg met glu asp

SEO ID No.45ZA (continued)

FIGURE 45ZA (continued)



fragment seq45ZA shifted minus 1 for the reading frame

1/1 31/11
ate tee eeg gac ace agg tea tee gge gag atg gtg ate gag get egg ace ege agg eat
ile ser pro asp thr arg ser ser gly glu met val ile glu ala arg thr arg arg his
61/21 91/31
con atta acc aga age acc age ate age aac ate geg atg gee age atg eeg ege egt egg
pro val ala arg gly thr ser ile ser asn ile ala met ala ser met pro arg arg
121/41
ate off ace act age gat cot tag gat gac age age age ace age toa
val leu ala thr arg asp pro trp asp asp gly gly ala AMB leu ala arg thr arg ser
191/61 211/71
teg tge cag ace ggg cat gee geg teg gea age tgt egg geg egg gtt aga geg gta geg
ser cys gln thr gly his ala ala ser ala ser cys arg ala arg val arg ala val ala
241/91
tgc gac cca gga tgg cga atg ctc ggg ggt cac cgg cga agt ggt agc cgc gga tga tgt
cys asp pro gly trp arg met leu gly gly his arg arg ser gly ser arg gly ora cys
301/101 331/111
egg tga age cea ace gge ggt aca ace gee acg eee gat tgt eet cae egt tgg tet eeg
arg OPA ser pro thr gly gly thr thr ala thr pro asp cys pro his arg trp ser pro
361/121
gtg tgg aga gca gga cgt tgt cct cgt cgc gac cgg cta gca gtc ggc ggg cca acg cct
val trp arg ala gly arg cys pro arg arg asp arg leu ala val gly gly pro thr pro
421/141 451/151
ccc cga ggc cac ggc ctt gag cgc ggg gaa gga tgt gca att cag tca act cga agt agc
pro arg gly his gly leu glu arg gly glu gly cys ala ile gln ser thr arg ser ser
481/161 511/171 511/171
tgg tca tca gtc ggg cga tcg cta ggc gcg gaa agc cgc tgc gtt gca agc cca gta cca
trp ser ser val gly arg ser leu gly ala glu ser arg cys val ala ser pro val pro
cct gct gtt gcc acc act ggc cgg gcg ccc cgg gat agc cgt acg cca ctc cga gca ttg pro ala val ala thr thr gly arg ala pro arg asp ser arg thr pro leu arg ala leu
601/201 631/211 gcg cgt tgc tca gtt cgg cgg ccg acg gca gcg ccg tgg tgt cgg cgg cct cgg cct gtt
ala arg cys ser val arg arg pro thr ala ala pro trp cys arg arg pro arg pro val
egg ctg ccg tta cct cga cgg ccg cga ccg cct gcc agc cgc gcc gcc gga tgt gct cca
arg leu pro leu pro arg arg pro arg pro pro ala ser arg ala ala gly cys ala pro
721/241
ggg aga the egg egg egg aag tet egg tee eec teg egt age gea teg egt ega eat aca
ala thr leu gly arg ala lys ser arg cys pro trp gly ser ala ser arg arg his thr
781/261 811/271
gog too agg cat cac cga ggc ggc gct cca tat cgc tgg gcg gca gat cga tga gga ata
pro ser gly his his arg gly gly ala pro tyr arg trp ala ala asp arg OPA gly ile
041/291
tog con acq cac gat atc ctc ctc atg tga tga acc gat gcg tgc ttg cgc acc agt acc
ser pro thr arg gly val leu leu met OPA OPA thr asp ala cys leu arg thr ser ile
001/301 931/311
granges and are get dag are are dec det aga egg age the tag egt and get at the get
gly gln ala asp glu ala ala arg ala giy arg gly leu AMB arg met ala var ser are
961/331
gag etc atc act acg acg ccg ccg aga tag aat cgc ccg cga acc agt ggt acg gcg cag
gln leu val ala ala pro pro gly AMB asn arg pro arg thr ser gly thr ala gli

SEQ ID No.45ZB

FIGURE 45ZB

WALLA TO CALE RELEVAN CREATED WHERE AN WA

1051/351 1021/341 att qac etc qta tea tet gag tta gtt gee ege gea atg gge ate ege gtg tta teg qta ile asp leu val ser ser glu leu val ala arg ala met gly ile arg val leu ser val 1111/371 1081/361 tta cgt gac agt ctg tcg gca agg ggc gca tgc cac tct ccg atc atg agc agc gga leu arg asp ser leu ser ala arg arg asp ala cys his ser pro ile met ser ser gly 1171/391 1141/381 tgc ttg acc aga tcg aga gcg ctc tct acg ccg aag atc cca agt tcg cat cga gtg tcc cys leu thr arg ser arg ala leu ser thr pro lys ile pro ser ser his arg val ser 1201/401 1231/411 gtg gcg ggg gct tcc gcg cac cga ccg cgc ggc gcc tgc agg gcg cgg cgt tgt tca val ala gly ala ser ala his arg pro arg gly gly ala cys arg ala arg arg cys ser 1291/431 tca tcq qtc tqq qqa tqt tqq ttt ccq qcq tqq cqt tca aaq aga cca tqa tcq gaa gtt ser ser val trp gly cys trp phe pro ala trp arg ser lys arg pro OPA ser glu val 1351/451 1321/441 tcc cga tac tca gcg ttt tcg gtt ttg tcg tga tgt tcg gtg gtg tgg tgt atg cca tca ser arg tyr ser ala phe ser val leu ser OPA cys ser val val trp cys met pro ser 1411/471 ccq qtc ctc ggt tgt ccg gca gga tgg atc gtg gcg gat cgg ctg ctg ggg ctt cgc gcc pro val leu gly cys pro ala gly trp ile val ala asp arg leu leu gly leu arg ala 1471/491 1441/481 age qte qta eea agg ggg eeg ggg get eat tea eea gee gta tgg aag ate ser val val pro arg gly pro gly ala his ser pro ala val trp lys ile

SEQ ID No.45ZB (continued)

FIGURE 45ZB (continued)

fragment seq45ZA shifted minus 2 for the reading frame

```
tet eee egg aca eea ggt cat eeg geg aga tgg tga teg agg ete gga eee gea gge ate
ser pro arg thr pro gly his pro ala arg trp OPA ser arg leu gly pro ala gly ile
61/21
                                        91/31
cgg tag cca gag gca cca gca tca gca aca tcg cga tgg cca gca tgc cgc gcc gtc qqq
arg AMB pro glu ala pro ala ser ala thr ser arg trp pro ala cys arg ala val gly
121/41
                                        151/51
tee ttg cea etc geg atc ett ggg atg acg gtg ggg eat age tag ege gea eea ggt eat
ser leu pro leu ala ile leu gly met thr val gly his ser AMB arg ala pro gly his
181/61
                                        211/71
cgt gcc aga ccg ggc atg ccg cgt cgg caa gct gtc ggg cgc ggg tta gag cgg tag cqt
arg ala arg pro gly met pro arg arg gln ala val gly arg gly leu glu arg AMB arg
                                        271/91
241/81
geg ace eag gat gge gaa tge teg ggg gte ace gge gaa gtg gta gee geg gat gat gte
ala thr gln asp gly glu cys ser gly val thr gly glu val val ala ala asp asp val
301/101
                                        331/111
ggt gaa gcc caa ccg gcg gta caa ccg cca cgc ccg att gtc ctc acc gtt ggt ctc cgg
gly glu ala gln pro ala val gln pro pro arg pro ile val leu thr val gly leu arg
361/121
                                        391/131
tgt gga gag cag gac gtt gtc ctc gtc gcg acc ggc tag cag tcg gcg qqc caa cqc ctc
cys gly glu gln asp val val leu val ala thr gly AMB gln ser ala gly gln arg leu
421/141
                                        451/151
ccc gag gcc acg gcc ttg agc gcg ggg aag gat gtg caa ttc agt caa ctc gaa gta gct
pro glu ala thr ala leu ser ala gly lys asp val gln phe ser gln leu glu val ala
481/161
                                        511/171
ggt cat cag tcg ggc gat cgc tag gcg cgg aaa gcc gct gcg ttg caa gcc cag tac cac
gly his gln ser gly asp arg AMB ala arg lys ala ala ala leu gln ala gln tyr his
541/181
                                        571/191
ctg ctg ttg cca cca ctg gcc ggg cgc ccc ggg ata gcc gta cgc cac tcc gag cat tgg
leu leu pro pro leu ala gly arg pro gly ile ala val arg his ser glu his trp
                                        631/211
ege gtt get eag tte gge gge ega egg eag ege egt ggt gte gge gge ete gge etq tte
arg val ala gln phe gly gly arg arg gln arg arg gly val gly gly leu gly leu phe
661/221
                                        691/231
gge tge egt tae etc gae gge ege ege etg eea gee geg eeg eeg gat gtg etc eag
gly cys arg tyr leu asp gly arg asp arg leu pro ala ala pro pro asp val leu gln
721/241
                                        751/251
cca cat tgg ggc gcg caa agt ctc ggt gcc cct ggg gta gcg cat cgc gtc gac ata cac
pro his trp gly ala gln ser leu gly ala pro gly val ala his arg val asp ile his
781/261
                                        811/271
cqt caq qqc atc acc gag gcg gcg ctc cat atc gct ggg cgg cag atc gat gag gaa tat
arg gln gly ile thr glu ala ala leu his ile ala gly arg gln ile asp glu glu tyr
                                        871/291
cgc caa cgc gcg gtg tcc tcc tca tgt gat gaa ccg atg cgt gct tgc gca cca gta tcg
arg gln arg ala val ser ser ser cys asp glu pro met arg ala cys ala pro val ser
901/301
                                        931/311
gac aag ccg atg agg ccg ccc gcg ctg gac ggg gct tgt agc gta tgg ccg ttt ccg ctc
asp lys pro met arg pro pro ala leu asp gly ala cys ser val trp pro phe pro leu
```

SEQ ID No.45ZC

FIGURE 45ZC

--- - - -



991/331 961/321 ago tog tog otg ogo ogo ogo gat aga ato goo ogo gaa oca gtg gta ogg ogo aga ser ser ser leu arg arg arg asp arg ile ala arg glu pro val val arg arg arg 1021/341 1051/351 ttg acc tcg tat cat ctg agt tag ttg ccc gcg caa tgg gca tcc gcg tgt tat cgg tat leu thr ser tyr his leu ser AMB leu pro ala gln trp ala ser ala cys tyr arg tyr 1111/371 tac gtg aca gtc tgt cgg caa gga ggg acg cat gcc act ctc cga tca tga gca gcg gat tyr val thr val cys arg gln gly gly thr his ala thr leu arg ser OPA ala ala asp 1171/391 1141/381 get tga eca gat ega gag ege tet eta ege ega aga tee eaa gtt ege ate gag tgt eeg ala OPA pro asp arg glu arg ser leu arg arg arg ser gln val arg ile glu cys pro 1231/411 1201/401 tgg egg ggg ett eeg ege ace gae ege geg geg eet gea ggg ege gge gtt gtt eat trp arg gly leu pro arg thr asp arg ala ala ala pro ala gly arg gly val val his 1291/431 1261/421 cat cgg tct ggg gat gtt ggt ttc cgg cgt ggc gtt caa aga gac cat gat cgg aag ttt his arg ser gly asp val gly phe arg gly val gln arg asp his asp arg lys phe 1351/451 1321/441 ccc gat act cag cgt ttt cgg ttt tgt cgt gat gtt cgg tgt gtg tgt ggt gta tgc cat cac pro asp thr gln arg phe arg phe cys arg asp val arg trp cys gly val cys his his 1411/471 1381/461 egg tee teg gtt gte egg eag gat gga teg tgg egg ate gge tge tgg gge tte geg eea arg ser ser val val arg gln asp gly ser trp arg ile gly cys trp gly phe ala pro 1471/491 gcg tcg tac caa ggg ggc cgg ggg ctc att cac cag ccg tat gga aga tc ala ser tyr gln gly gly arg gly leu ile his gln pro tyr gly arg

SEQ ID No.45ZC (continued 1)

FIGURE 45ZC (continued 1)

ORF de seq 45ZA directement en fusion avec phoA cag tot gtc ggc aag gag gga cgc atg cca ctc tcc gat cat gag cag cgg gln ser val gly lys glu gly arg met pro leu ser asp his glu gln arg 1171/391 1141/381 atg ctt gac cag atc gag agc gct ctc tac gcc gaa gat ccc aag ttc gca tcg agt gtc met leu asp gln ile glu ser ala leu tyr ala glu asp pro lys phe ala ser ser val 1201/401 1231/411 cgt ggc ggg ggc ttc cgc gca ccg acc gcg cgg cgc ctg cag ggc gcg gcg ttg ttc arg gly gly phe arg ala pro thr ala arg arg leu gln gly ala ala leu phe 1291/431 atc atc ggt ctg ggg atg ttg gtt tcc ggc gtg gcg ttc aaa gag acc atg atc gga agt ile ile gly leu gly met leu val ser gly val ala phe lys glu thr met ile gly ser 1351/451 tto cog ata oto ago gtt tto ggt ttt gto gtg atg tto ggt ggt gtg tat god ato phe pro ile leu ser val phe gly phe val val met phe gly gly val val tyr ala ile 1411/471 ace gqt cet egg ttg tee gge agg atg gat egt gge gga teg get get ggg get teg ege thr gly pro arg leu ser gly arg met asp arg gly gly ser ala ala gly ala ser arg 1441/481 1471/491 cag cgt cgt acc aag ggg gcc ggg ggc tca ttc acc agc cgt atg gaa gat c gln arg arg thr lys gly ala gly gly ser phe thr ser arg met glu asp

SEQ ID No.45A

FIGURE 45A

Sequence Rv2169c predicted by Cole et al., 1998 (Nature 393:537-544) and containing Seq45A 1/1 31/11 atg cca ctc tcc gat cat gag cag cgg atg ctt gac cag atc gag agc gct ctc tac gcc Met pro leu ser asp his glu gln arg met leu asp gln ile glu ser ala leu tyr ala 91/31 61/21 gaa gat ccc aag ttc gca tcg agt gtc cgt ggc ggg ggc ttc cgc gca ccg acc gcg cgg glu asp pro lys phe ala ser ser val arg gly gly phe arg ala pro thr ala arg 151/51 egg ege etg eag gge geg ttg tte ate ate ggt etg ggg atg ttg gtt tee gge gtg arg arg leu gln gly ala ala leu phe ile ile gly leu gly met leu val ser gly val 211/71 181/61 gcg ttc aaa gag acc atg atc gga agt ttc ccg ata ctc agc gtt ttc ggt ttt gtc gtg ala phe lys glu thr met ile gly ser phe pro ile leu ser val phe gly phe val val 271/91 241/81 atg ttc ggt ggt gtg tat gcc atc acc ggt cct cgg ttg tcc ggc agg atg gat cgt met phe gly gly val val tyr ala ile thr gly pro arg leu ser gly arg met asp arg 331/111 301/101 gge gga teg get get ggg get teg ege eag egt egt ace aag ggg gee ggg gge tea tte gly gly ser ala ala gly ala ser arg gln arg arg thr lys gly ala gly gly ser phe 391/131 361/121 acc agc cgt atg gaa gat cgg ttc cgg cgc cgc ttc gac gag taa thr ser arg met glu asp arg phe arg arg phe asp glu OCH

SEQ ID No.45D

FIGURE 45D

ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv2169c

31/11 tga cag tot gto ggo aag gag gga ogo atg coa oto too gat cat gag cag ogg atg ott OPA gln ser val gly lys glu gly arg met pro leu ser asp his glu gln arg met leu 91/31 gac cag atc gag agc gct ctc tac gcc gaa gat ccc aag ttc gca tcg agt gtc cgt ggc asp gln ile glu ser ala leu tyr ala glu asp pro lys phe ala ser ser val arg gly 151/51 qqq qqc ttc cgc gca ccg acc gcg cgg cgc ctg cag ggc gcg gcg ttg ttc atc atc gly gly phe arg ala pro thr ala arg arg leu gln gly ala ala leu phe ile ile 211/71 qgt ctg ggg atg ttg gtt tcc ggc gtg gcg ttc aaa gag acc atg atc gga agt ttc ccg gly leu gly met leu val ser gly val ala phe lys glu thr met ile gly ser phe pro 271/91 ata etc age gtt tte ggt ttt gte gtg atg tte ggt gtg gtg tat gee ate ace ggt ile leu ser val phe gly phe val val met phe gly gly val val tyr ala ile thr gly 331/111 cct cgg ttg tcc ggc agg atg gat cgt ggc gga tcg gct gct ggg gct tcg cgc cag cgt pro arg leu ser gly arg met asp arg gly gly ser ala ala gly ala ser arg gln arg 361/121 391/131 cgt acc aag ggg gcc ggg ggc tca ttc acc agc cgt atg gaa gat cgg ttc cgg cgc cgc arg thr lys gly ala gly gly ser phe thr ser arg met glu asp arg phe arg arg arg 421/141 ttc gac gag taa phe asp glu OCH

SEQ ID No.45F

FIGURE 45F

1/1 31/11 cag ccg cgc cgc atc gac cag ggc ctc acg ccc ggt cac ttc tcc gcg ttc ctc aac aat gln pro arg arg ile asp gln gly leu thr pro gly his phe ser ala phe leu asn asn 61/21 91/31 tec ggt gaa cat ege acc agg tta gge age aat eee geg gae eeg cac eee act ege ega ser gly glu his arg thr arg leu gly ser asn pro ala asp pro his pro thr arg arg 151/51 121/41 ccg gcc aac tca cag aca ccc tct acg atg cag ggt atg cgg acc ccc aga cgc cac tgc pro ala asn ser gln thr pro ser thr met gln gly met arg thr pro arg arg his cys 181/61 211/71 cgt cgc atc gcc gtc ctc gcc gcc gtt agc atc gcc gcc act gtc gtt gcc ggc tgc tcg arg arg ile ala val leu ala ala val ser ile ala ala thr val val ala gly cys ser 271/91 241/81 teg gge teg aag eea age gge gga eea ett eeg gae geg aag eeg etg gte gag gae gee ser gly ser lys pro ser gly gly pro leu pro asp ala lys pro leu val glu glu ala 301/101 331/111 acc gcg cag acc aag gct ctc aag agc gcg cac atg gtg ctg acg gtc aac ggc aag atc thr ala gln thr lys ala leu lys ser ala his met val leu thr val asn gly lys ile

SEQ ID No.46A

FIGURE 46A

1/1									31/1	.1								
agc cgc	gcc	gca	tcg	acc	agg	gcc	tca	cgc	ccg	gtc	act	tct	ccg	cgt	tcc	tca	aca	att
ser arg	ala	ala	ser	thr	arg	ala	ser	arg	pro	val	thr	ser	pro	arg	ser	ser	thr	ile
61/21									91/3	31								
ccg gtg	aac	atc	gca	cca	ggt	tag	gca	gca	atc	ccg	cgg	acc	cgc	acc	cca	ctc	gcc	gac
pro val	asn	ile	ala	pro	gly	AMB	ala	ala	ile	pro	arg	thr	arg	thr	pro	leu	ala	asp
121/41									151/	'51								
cgg cca	act	cac	aga	cac	cct	cta	cga	tgc	agg	gta	tgc	gga	CCC	cca	gac	gcc	act	gcc
arg pro	thr	his	arg	his	pro	leu	arg	cys	arg	val	cys	gly	pro	pro	asp	ala	thr	ala
181/61									211/	71								
101/01									/									
gtc gca	tcg	ccg	tcc	tcg	ccg	ccg	tta	gca	,		cca	ctg	tcg	ttg	ccg	gct	gct	cgt
•	_	_		_	_	-		-	tcg	ccg			_	_	-	-	-	-
gtc gca	_	_		_	_	-		-	tcg	ccg pro			_	_	-	-	-	-
gtc gca val ala	ser	pro	ser	ser	pro	pro	leu	ala	tcg ser 271/	ccg pro '91	pro	leu	ser	leu	pro	ala	ala	arg
gtc gca val ala 241/81	ser cga	pro agc	ser caa	ser	pro	pro gac	leu cac	ala	tcg ser 271/ cgg	ccg pro '91 acg	pro cga	leu agc	ser	leu tgg	pro tcg	ala agg	ala agg	arg
gtc gca val ala 241/81 cgg gct	ser cga	pro agc	ser caa	ser	pro	pro gac	leu cac	ala	tcg ser 271/ cgg	ccg pro '91 acg thr	pro cga	leu agc	ser	leu tgg	pro tcg	ala agg	ala agg	arg
gtc gca val ala 241/81 cgg gct arg ala	ser cga arg	pro agc ser	ser caa gln	ser gcg ala	pro gcg ala	pro gac asp	leu cac his	ala ttc phe	tcg ser 271/ cgg arg 331/	ccg pro '91 acg thr	pro cga arg	leu agc ser	ser cgc arg	leu tgg trp	pro tcg ser	ala agg arg	ala agg arg	arg cca pro

SEQ ID No.46B

FIGURE 46B



1/1										31/1	1								
gcc	gcg	ccg	cat	cga	cca	ggg	cct	cac	gcc	cgg	tca	ctt	ctc	cgc	gtt	cct	caa	caa	ttc
ala	ala	pro	his	arg	pro	gly	pro	his	ala	arg	ser	leu	leu	arg	val	pro	gln	gln	phe
61/2	21									91/3	31								
		aca																	
arg	OPA	thr	ser	his	gln	val	arg	gln	gln	ser	arg	gly	pro	ala	pro	his	ser	pro	thr
121/	41									151/	/51								
		ctc																	
gly	gln	leu	thr	asp	thr	leu	tyr	asp	ala	gly	tyr	ala	asp	pro	gln	thr	pro	leu	pro
181/	61									211/	71								
tcg	cat	cgc	cgt	cct	cgc	cgc	cgt	tag	cat	cgc	cgc	cac	tgt	cgt	tgc	cgg	ctg	ctc	gtc
ser	his	arg	arg	pro	arg	arg	arg	AMB	his	arg	arg	his	cys	arg	cys	arg	leu	leu	val
241/	81									271/	91								
ggg	ctc	gaa	gcc	aag	cgg	cgg	acc	act	tcc	gga	cgc	gaa	gcc	gct	ggt	cga	gga	ggc	cac
gly	leu	glu	ala	lys	arg	arg	thr	thr	ser	gly	arg	glu	ala	ala	gly	arg	gly	gly	\mathtt{his}
301/	101									331,	1111								
cgc	gca	gac	caa	ggc	tct	caa	gag	cgc	gca	cat	ggt	gct	gac	ggt	caa	cgg	caa	gat	С
arg	ala	asp	gln	gly	ser	gln	glu	arg	ala	his	gly	ala	asp	gly	gln	arg	gln	asp	

SEQ ID No.46C

FIGURE 46C



Coding sequence Rv1411c predicted by Cole et al., 1998 (Nature 393: 537-544) and containing seq46A:

```
31/11
1/1
atg egg acc eec aga ege cac tge egt ege ate gee gte ete gee gee gtt age ate gee
Met arg thr pro arg arg his cys arg arg ile ala val leu ala ala val ser ile ala
                                        91/31
ged act gtd gtd gcd ggd tgg tcg tcg ggd cca agd ggd gga cca ctt ccg gad
ala thr val val ala gly cys ser ser gly ser lys pro ser gly gly pro leu pro asp
                                        151/51
121/41
gcg aag ccg ctg gtc gag gag gcc acc gcg cag acc aag gct ctc aag agc gcg cac atg
ala lys pro leu val glu glu ala thr ala gln thr lys ala leu lys ser ala his met
181/61
                                        211/71
qtq ctq acq qtc aac ggc aag atc ccg gga ctg tct ctg aag acg ctg agc ggc gat ctc
val leu thr val asn gly lys ile pro gly leu ser leu lys thr leu ser gly asp leu
                                        271/91
241/81
ace ace ace eec ace gee geg acg gga aac gte aag etc acg etg ggt ggg tet gat ate
thr thr asn pro thr ala ala thr gly asn val lys leu thr leu gly gly ser asp ile
                                        331/111
qat qcc gac ttc gtg gtg ttc gac ggg atc ctg tac gcc acc ctg acg ccc aac cag tgg
asp ala asp phe val val phe asp gly ile leu tyr ala thr leu thr pro asn gln trp
                                        391/131
361/121
age gat the ggt eec gee gae ate tae gae eec gee eag gtg etg aat eeg gat ace
ser asp phe gly pro ala ala asp ile tyr asp pro ala gln val leu asn pro asp thr
                                        451/151
421/141
qgc ctg qcc aac gtg ctg gcg aat ttc gcc gac gca aaa gcc gaa ggg cgg gat acc atc
gly leu ala asn val leu ala asn phe ala asp ala lys ala glu gly arg asp thr ile
                                        511/171
481/161
aac qgc cag aac acc atc cgc atc agc ggg aag gta tcg gca cag gcg gtg aac cag ata
asn gly gln asn thr ile arg ile ser gly lys val ser ala gln ala val asn gln ile
                                        571/191
qcq ccq ccq ttc aac gcg acg cag ccg gtg ccg gcg acc gtc tgg att cag gag acc ggc
ala pro pro phe asn ala thr gln pro val pro ala thr val trp ile gln glu thr gly
                                        631/211
601/201
gat cat caa ctg gca cag gcc cag ttg gac cgc ggc tcg ggc aat tcc gtc cag atg acc
asp his gln leu ala gln ala gln leu asp arg gly ser gly asn ser val gln met thr
                                        691/231
661/221
ttg tcg aaa tgg ggc gag aag gtc cag gtc acg aag ccc ccg gtg agc tga
leu ser lys trp gly glu lys val gln val thr lys pro pro val ser OPA
```

SEQ ID No.46D

FIGURE 46D



ORF according to Cole et al., 1998 (Nature 393: 537-544): and containing the coding sequence Rv1411c:

31/11 tag etc ace eag gtt gga eeg gtt eag tgt etc gge eat eac gte gge ggt gaa ttg gee AMB leu thr gln val gly pro val gln cys leu gly his his val gly gly glu leu ala 91/31 61/21 gtc ggg caa tac atc gac ggc cgt cag aca cac gcc gtt gac agc gat cga gtc gcc gtg val gly gln tyr ile asp asp arg gln thr his ala val asp ser asp arg val ala val 151/51 qcc qqc qtc qgc ggt aac cat cgg acc gcg gat ggt cag ccg cgc cgc atc gac caq ggc ala gly val gly gly asn his arg thr ala asp gly gln pro arg arg ile asp gln gly 181/61 211/71 ctc acg ccc ggt cac ttc tcc gcg ttc ctc aac aat tcc ggt gaa cat cgc acc agg tta leu thr pro gly his phe ser ala phe leu asn asn ser gly glu his arg thr arg leu 241/81 271/91 gge age aat eee geg gae eeg cac eee act ege ega eeg gee aac tea eag aca eee tet gly ser asn pro ala asp pro his pro thr arg arg pro ala asn ser gln thr pro ser 331/111 301/101 acg atg cag ggt atg cgg acc ccc aga cgc cac tgc cgt cgc atc gcc gtc ctc gcc gcc thr met gln gly met arg thr pro arg arg his cys arg arg ile ala val leu ala ala 391/131 gtt age ate gee gee act gte gtt gee gge tge teg teg gge teg aag eea age gge gga val ser ile ala ala thr val val ala gly cys ser ser gly ser lys pro ser gly gly 451/151 421/141 cca ctt ccg gac gcg aag ccg ctg gtc gag gag gcc acc gcg cag acc aag gct ctc aag pro leu pro asp ala lys pro leu val glu glu ala thr ala gln thr lys ala leu lys 511/171 481/161 ago gog cao atg gtg ctg acg gtc aac ggc aag atc ccg gga ctg tct ctg aag acg ctg ser ala his met val leu thr val asn gly lys ile pro gly leu ser leu lys thr leu 571/191 541/181 age gge gat etc ace ace aac eec ace gee geg aeg gga aac gte aag etc aeg etg ggt ser gly asp leu thr thr asn pro thr ala ala thr gly asn val lys leu thr leu gly 601/201 631/211 qqq tot gat ato gat goo gad tto gtg gtg tto gad ggg ato otg tad god acd otg acq gly ser asp ile asp ala asp phe val val phe asp gly ile leu tyr ala thr leu thr 691/231 661/221 ccc aac caq tgg age gat tte ggt eee gee gee gae ate tae gae eee gee eag gtg etg pro asn gln trp ser asp phe gly pro ala ala asp ile tyr asp pro ala gln val leu 751/251 721/241 aat eeg gat ace gge etg gee aac gtg etg geg aat tte gee gae gea aaa gee gaa ggg asn pro asp thr gly leu ala asn val leu ala asn phe ala asp ala lys ala glu gly 781/261 811/271 cgg gat acc atc aac ggc cag aac acc atc cgc atc agc ggg aag gta tcg gca cag gcg arg asp thr ile asn gly gln asn thr ile arg ile ser gly lys val ser ala gln ala 871/291 841/281 qtq aac caq ata qcq ccq ccq ttc aac qcq acq caq ccq gtq ccq gcq acc gtc tgg att val asn qln ile ala pro pro phe asn ala thr gln pro val pro ala thr val trp ile 931/311 cag gag acc ggc gat cat caa ctg gca cag gcc cag ttg gac cgc ggc tcg ggc aat tcc gln glu thr gly asp his gln leu ala gln ala gln leu asp arg gly ser gly asn ser 991/331 gtc cag atg acc ttg tcg aaa tgg ggc gag aag gtc cag gtc acg aag ccc ccg gtg agc val gln met thr leu ser lys trp gly glu lys val gln val thr lys pro pro val ser 1021/341 tga OPA

SEQ ID No.46F



31/11 1/1qaq ctq qtc aac qqc gcc ggc atc gac gcc gcc gtc gtg acc tgc cgg ccg gac agc glu leu val asn gly ala gly ile asp asp ala ala val val thr cys arg pro asp ser 91/31 61/21 ctg gec gat gec cag cag atg gtc gag geg gea etg ggc ega tat ggc egt ttg gac gga leu ala asp ala gln gln met val glu ala ala leu gly arg tyr gly arg leu asp gly 151/51 gtg ttg gtg gcc tcg ggc agc aac cat gtg gcg ccc att acc gag atg gcc gtc gag gac val leu val ala ser gly ser asn his val ala pro ile thr glu met ala val glu asp 211/71 181/61 phe asp ala val met asp ala asn val arg gly ala trp leu val cys arg ala ala gly 271/91 cgg gtg ctg ctc gag cag ggt cag ggc ggc agc gtg gtg ctg gtg tcg tcc gtt cgc ggc arg val leu leu glu gln gly gln gly gly ser val val leu val ser ser val arg gly 331/111 301/101 ggg ttg ggc aat gcc gcc ggt tac agc gcg tac tgc ccg tcg aag gcg ggc acc gat c gly leu gly asn ala ala gly tyr ser ala tyr cys pro ser lys ala gly thr asp

SEQ ID No.47A

FIGURE 47A

agc tgg tca acg gcg ccg gca tcg acg acg ccg ccg tcg tcg tga cct gcc ggc cgg aca gcc ser trp ser thr ala pro ala ser thr thr pro pro ser OPA pro ala gly arg thr ala 61/21 tgg ccg atg ccc agc aga tgg tcg agg cgg cac tgg gcc gat atg gcc gtt tgg acg gag trp pro met pro ser arg trp ser arg arg his trp ala asp met ala val trp thr glu 121/41 tgt tgg tgg cct cgg gca gca acc atg tgg cgc cca tta ccg aga tgg ccg tcg agg act cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt tgt gtc ggg cgg c	1/1										31/1	1								
tgg ccg atg ccc agc aga tgg tcg agg cgg cac tgg gcc gat atg gcc gtt tgg acg gag trp pro met pro ser arg trp ser arg arg his trp ala asp met ala val trp thr glu 151/41 tgt tgg tgg cct cgg gca gca acc atg tgg cgc cca tta ccg aga tgg ccg tcg agg act cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	agc	tgg	tca	acg	gcg	ccg	gca	tcg	acg	acg	ccg	ccg	tcg	tga	cct	gcc	ggc	cgg	aca	gcc
tgg ccg atg ccc agc aga tgg tcg agg cgg cac tgg gcc gat atg gcc gtt tgg acg gag trp pro met pro ser arg trp ser arg arg his trp ala asp met ala val trp thr glu 121/41 tgt tgg tgg cct cgg gca gca acc atg tgg cgc cca tta ccg aga tgg ccg tcg agg act cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	ser	trp	ser	thr	ala	pro	ala	ser	thr	thr	pro	pro	ser	OPA	pro	ala	gly	arg	thr	ala
trp pro met pro ser arg trp ser arg arg his trp ala asp met ala val trp thr glu 121/41	61/2	1									91/3	31								
121/41 tgt tgg tgg cct cgg gca gca acc atg tgg cgc cca tta ccg aga tgg ccg tcg agg act cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	tgg	ccg	atg	ccc	agc	aga	tgg	tcg	agg	cgg	cac	tgg	gcc	gat	atg	gcc	gtt	tgg	acg	gag
121/41 tgt tgg tgg cct cgg gca gca acc atg tgg cgc cca tta ccg aga tgg ccg tcg agg act cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	trp	pro	met	pro	ser	arg	trp	ser	arg	arg	his	trp	ala	asp	met	ala	val	trp	thr	glu
cys trp trp pro arg ala ala thr met trp arg pro leu pro arg trp pro ser arg thr 181/61 211/71 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c																				
181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	tgt	tgg	tgg	cct	cgg	gca	gca	acc	atg	tgg	cgc	cca	tta	ccg	aga	tgg	ccg	tcg	agg	act
181/61 tcg acg ctg tga tgg acg cga acg tgc ggg gtg cct ggc tgg tgt gtc ggg cgg c	cys	trp	trp	pro	arg	ala	ala	thr	met	trp	arg	pro	leu	pro	arg	trp	pro	ser	arg	thr
ser thr leu OPA trp thr arg thr cys gly val pro gly trp cys val gly arg pro asp 241/81 271/91 ggg tgc tgc tcg agc agg gtc agg gcg gca gcg tgg tgc tgg tgt cgt ccg ttc gcg gcg			_	_																
241/81 ggg tgc tgc tcg agc agg gtc agg gcg gca gcg tgg tgc tgg tgt cgt ccg ttc gcg gcg	tcg	acg	ctg	tga	tgg	acg	cga	acg	tgc	ggg	gtg	cct	ggc	tgg	tgt	gtc	ggg	cgg	ccg	gac
ggg tgc tgc tcg agc agg gtc agg gcg gca gcg tgg tgc tgg tgt cgt ccg ttc gcg gcg	ser	thr	leu	OPA	trp	thr	arg	thr	cys	gly	val	pro	gly	trp	cys	val	gly	arg	pro	asp
gly cys cys ser ser arg val arg ala ala ala trp cys trp cys arg pro phe ala ala 301/101 331/111 ggt tgg gca atg ccg ccg gtt aca gcg cgt act gcc cgt cga agg cgg gca ccg atc	241/	81									271	/91								
301/101 331/111 ggt tgg gca atg ccg ccg gtt aca gcg cgt act gcc cgt cga agg cgg gca ccg atc	ggg	tgc	tgc	tcg	agc	agg	gtc	agg	gcg	gca	gcg	tgg	tgc	tgg	tgt	cgt	ccg	ttc	gcg	gcg
ggt tgg gca atg ccg ccg gtt aca gcg cgt act gcc cgt cga agg cgg gca ccg atc	gly	cys	cys	ser	ser	arg	val	arg	ala	ala	ala	trp	cys	trp	cys	arg	pro	phe	ala	ala
	301/	101									331,	/111								
	ggt	tgg	gca	atg	ccg	ccg	gtt	aca	gcg	cgt	act	gcc	cgt	cga	agg	cgg	gca	ccg	atc	
gly trp ala met pro pro val thr ala arg thr ala arg arg arg ala pro ile	gly	trp	ala	met	pro	pro	val	thr	ala	arg	thr	ala	arg	arg	arg	arg	ala	pro	ile	

SEQ ID No.47B

FIGURE 47B



1/1									31/1	1								
gct ggt	caa	cgg	cgc	cgg	cat	cga	cga	cgc	cgc	cgt	cgt	gac	ctg	ccg	gcc	gga	cag	cct
ala gly	gln	arg	arg	arg	his	arg	arg	arg	arg	arg	arg	asp	leu	pro	ala	gly	gln	pro
61/21									91/3	31								
ggc cga	tgc	cca	gca	gat	ggt	cga	ggc	ggc	act	ggg	ccg	ata	tgg	ccg	ttt	gga	cgg	agt
gly arg	cys	pro	ala	asp	gly	arg	gly	gly	thr	gly	pro	ile	trp	pro	phe	gly	arg	ser
121/41									151,	/51								
gtt ggt	ggc	ctc	ggg	cag	caa	cca	tgt	ggc	gcc	cat	tac	cga	gat	ggc	cgt	cga	gga	ctt
val gly	gly	leu	gly	gln	gln	pro	cys	gly	ala	his	tyr	arg	asp	gly	arg	arg	дſУ	leu
181/61									211,	/71								
cga cgc																		
arg arg	cys	asp	gly	arg	glu	arg	ala	gly	cys	leu	ala	gly	val	ser	gly	gly	arg	thr
241/81									271	/91								
ggt gct																		
gly ala	ala	arg	ala	gly	ser	gly	arg	gln	arg	gly	ala	gly	val	val	arg	ser	arg	arg
301/101									331,	/111								
gtt ggg	caa	tgc	cgc	cgg	tta	cag	cgc	gta	ctg	CCC	gtc	gaa	ggc	ggg	cac	cga	tc	
val gly	gln	cys	arg	arg	leu	gln	arg	val	leu	pro	val	glu	gly	gly	his	arg		

SEQ ID No.47C

FIGURE 47C



Coding sequence Rv1714 predicted by Cole et al., 1998 (Nature 393: 537-544) and containing seq 47A:

1/1	31/11
ata ana dan ata aca cta act cag cag ata	ccg aac ctg ggt ctg gcg cgc ttc agc gtg
val glu glu met ala leu ala gln gln val	pro asn leu gly leu ala arg phe ser val
61/21	91/31
and was san ten ate eth ate ace ddc dcd	acc ggt tcg ttg ggc cga gtt gcc gcc cgg
gln asp lys ser ile leu ile thr gly ala	thr gly ser leu gly arg val ala ala arg
121/41	151/51
gcg ctg gcc gac gcg gga gcg cgg ctg aca	ctg gcc ggc ggc aac tcg gcc ggt ctg gcc
ala leu ala asp ala gly ala arg leu thr	leu ala gly gly asn ser ala gly leu ala 211/71
181/61	
gag ctg gtc aac ggc gcc ggc atc gac gac	gcc gcc gtc gtg acc tgc cgg ccg gac agc
	ala ala val val thr cys arg pro asp ser 271/91
241/81	gca ctg ggc cga tat ggc cgt ttg gac gga
ctg gcc gat gcc cag cag acg gcc gay gcg	ala leu gly arg tyr gly arg leu asp gly
	331/111
301/101	gcg ccc att acc gag atg gcc gtc gag gac
gtg ttg gtg gtc teg ggc age aac out gog	ala pro ile thr glu met ala val glu asp
261/121	391/131
tte mag get gtg atg gac gcg aac gtg cgc	ggt gcc tgg ctg gtg tgt cgg gcg gcc gga
phe asp ala val met asp ala asn val arg	gly ala trp leu val cys arg ala ala gly
121/111	451/151
and ata ota oto dad cad dat cad dac dag	age gtg gtg ctg gtg tcg tcc gtt cgc ggc
arg val leu leu glu gln gly gln gly gly	ser val val leu val ser ser val arg gry
401 /161	511/1/1
ggg ttg ggc aat gcc gcc ggt tac agc gcc	g tac tgc ccg tcg aag gcg ggc acc gat ctg
gly leu gly asn ala ala gly tyr ser ala	a tyr cys pro ser lys ala gly thr asp leu
541/181	571/191
ttg gcc aag aca ttg gcg gcc gaa tgg gg	ggt cac ggc att cgg gtg aac gcg ctg gcg
	y gly his gly ile arg val asn ala leu ala 631/211
601/201	g tgg atg ttc acc gac gat ccg aag ggc cgg
ccg acg gtg ttt cgg tcc gcg gtg acc ga	u trp met phe thr asp asp pro lys gly arg
	691/231
661/221	c ccg ttg cgc cgc ttc gcc gaa ccg gaa gac
gcc acc egg gag geg acg eee gee egg ac	e pro leu arg arg phe ala glu pro glu asp
721/241	751/251
the ste see see ste ato tat cte ste ag	c gac gcc tcg agc ttc tac acc ggc cag gtg
nhe val glv ala leu ile tyr leu leu se	r asp ala ser ser phe tyr thr gly gln val
781/261	811/2/1
atg tat ctg gac ggc ggg tac acc gca tg	c tga
met tyr leu asp gly gly tyr thr ala cy	s OPA

SEQ ID No.47D

FIGURE 47D

THE TANKERS OF THE PARTY OF THE



ORF according to Cole et al., 1998 (Nature 393: 537-544) and containing the coding sequence Rv1714:

04/1	54/11
24/1	gtg ccg aac ctg ggt ctg gcg cgc ttc agc
AMP wal glu glu met ala leu ala gln gln	val pro asn leu gly leu ala arg phe ser
04/21	114/31 .
The gam gam and the ate etg ate acc gge	gcg acc ggt tcg ttg ggc cga gtt gcc gcc
val gln asp lys ser ile leu ile thr gly	ala thr gly ser leu gly arg val ala ala
144/47	174/51
and aca ota acc asc aca aga aca caa cta	aca ctg gcc ggc ggc aac tcg gcc ggt ctg
arg ala leu ala asp ala gly ala arg leu	thr leu ala gly gly ash ser ala gly leu
204/61	234/71
gcc gag ctg gtc aac ggc gcc ggc atc gac	gac gcc gcc gtc gtg acc tgc cgg ccg gac
ala glu leu val asn gly ala gly ile asp	asp ala ala val val thr cys arg pro asp
264/81	294/91
age etg gee gat gee eag eag atg gte gag	geg gea etg gge ega tat gge egt ttg gae
	ala ala leu gly arg tyr gly arg leu asp 354/111
324/101	gtg gcg ccc att acc gag atg gcc gtc gag
gga gtg ttg gtg gcc tcg ggc agc aac cac	s val ala pro ile thr glu met ala val glu
	414/131
384/121	g cgg ggt gcc tgg ctg gtg tgt cgg gcg gc
gac ttc gac gct gtg atg gac gcg aac gcg	arg gly ala trp leu val cys arg ala ala
	474/151
444/141	ggc agc gtg gtg ctg gtg tcg tcc gtt cgc
gga cgg gcg ccg ccc gug cug gly gly gly gly gly gly gly gly gly gl	y gly ser val val leu val ser ser val arg
E04/361	534/1/1
and the end sat end end ent tac age	geg tac tgc ccg tcg aag gcg ggc acc gat
gly gly leu gly asn ala ala gly tyr se	r ala tyr cys pro ser lys ala gly thr asp
ECA/101	594/191
ctg ttg gcc aag aca ttg gcg gcc gaa tg	g ggc ggt cac ggc att cgg gtg aac gcg ctg
leu leu ala lys thr leu ala ala glu tr	b did did us did the and has asu and ten
624/201	654/211
gcg ccg acg gtg ttt cgg tcc gcg gtg ac	c gag tgg atg ttc acc gac gat ccg aag ggc
ala pro thr val phe arg ser ala val th	r glu trp met pne thi asp asp pro 193 gry
684/221	714/231
cgg gcc acc cgg gag gcg atg ctc gcc cg	g atc ccg ttg cgc cgc ttc gcc gaa ccg gaa
	g ile pro leu arg arg phe ala glu pro glu 774/251
744/241	c age gac gec teg age tte tac ace ggc cag
gac ttc gtc ggc gcc ctg atc tat ctg cc	u ser asp ala ser ser phe tyr thr gly gln
	834/271
804/261	
gtg atg tat ctg gac ggc ggg tac acc gc	a cvs OPA
val met tyr leu asp gly gly tyr thr al	1

SEQ ID No.47F

FIGURE 47F

THE RESIDENCE OF THE PROPERTY OF THE PROPERTY

1/1 agg ctc atg agc aag acg gtt ctc atc ctt ggc gcg ggt gtc ggc ggc ctg acc acc gcc arg leu met ser lys thr val leu ile leu gly ala gly val gly gly leu thr thr ala 61/21 91/31 gac acc ctc cgt caa ctg cta cca cct gag gat c asp thr leu arg gln leu leu pro pro glu asp

SEQ ID No.48A

FIGURE 48A

1/1 ggc tca tga gca aga cgg ttc tca tcc ttg gcg cgg gtg tcg gcg gcc tga cca ccg ccg gly ser OPA ala arg arg phe ser ser leu ala arg val ser ala ala OPA pro pro pro 61/21 91/31 aca ccc tcc gtc aac tgc tac cac ctg agg atc thr pro ser val asn cys tyr his leu arg ile

SEQ ID No.48B

FIGURE 48B

1/1
gct cat gag caa gac ggt tct cat cct tgg cgc ggg tgt cgg cgg cct gac cac cgc cga
ala his glu gln asp gly ser his pro trp arg gly cys arg arg pro asp his arg arg
61/21
cac cct ccg tca act gct acc acc tga gga tc
his pro pro ser thr ala thr thr OPA gly

SEQ ID No.48C

FIGURE 48C

AN IT TITAL MICHELY MICHAELERS . THE



Coding sequence Rv0331 predicted by Cole et al., 1998 (Nature 393: 537-544) and containing seq48A:

```
31/11
atg age aag acg gtt ctc atc ctt ggc gcg ggt gtc ggc ggc ctg acc acc gcc gac acc
Met ser lys thr val leu ile leu gly ala gly val gly gly leu thr thr ala asp thr
                                        91/31
61/21
ctc cgt caa ctg cta cca cct gag gat cga atc ata ttg gtg gac agg agc ttt gac ggg
leu arg gln leu leu pro pro glu asp arg ile ile leu val asp arg ser phe asp gly
                                        151/51
121/41
acg ctg ggc ttg tcg ttg cta tgg gtg ttg cgg ggc tgg cgg cct gac gac gtc cgc
thr leu gly leu ser leu leu trp val leu arg gly trp arg arg pro asp asp val arg
                                        211/71
gtc ege eec ace geg geg teg etg eec ggt gtg gaa atg gtt act gea ace gte gee eac
val arg pro thr ala ala ser leu pro gly val glu met val thr ala thr val ala his
241/81
                                        271/91
att gac atc gcg gcc cag gta gtg cac acc gac aac agc gtc atc ggc tat gac gcg ttg
ile asp ile ala ala gln val val his thr asp asn ser val ile gly tyr asp ala leu
                                        331/111
301/101
gtg atc gca tta ggt gcg gcg ctg aac acc gac gcc gtt ccc gga ctg tcg gac gcg ctc
val ile ala leu gly ala ala leu asn thr asp ala val pro gly leu ser asp ala leu
                                        391/131
361/121
qac qcc qac qtc qcg qgc cag ttc tac acc ctg gac ggc gcg gct gag ctg cgt gcg aaq
asp ala asp val ala gly gln phe tyr thr leu asp gly ala ala glu leu arg ala lys
                                        451/151
421/141
gtc gag gcg ctc gag cat ggc cgg atc gct gtg gct atc gcc ggg gtg ccg ttc aaa tgc
val glu ala leu glu his gly arg ile ala val ala ile ala gly val pro phe lys cys
                                        511/171
481/161
cca gcc gca ccg ttc gaa gcg gcg ttt ctg atc gcc gcc caa ctc ggt gac cgc tac gcc
pro ala ala pro phe glu ala ala phe leu ile ala ala gln leu gly asp arg tyr ala
                                        571/191
acc gga acc gta cag atc gac acg ttc acg cet gac ccg ctg ccg atg ccc gtt gca ggt
thr gly thr val gln ile asp thr phe thr pro asp pro leu pro met pro val ala gly
                                        631/211
601/201
ccc gag gtc ggc gag gct ttg gtc tcg atg ctc aag gat cac ggt gtc ggc ttc cat cct
pro glu val gly glu ala leu val ser met leu lys asp his gly val gly phe his pro
661/221
                                        691/231
cgc aag gcc cta gct cgc gtc gat gag gcc gca agg acg atg cac ttc ggt gac ggc acg
arg lys ala leu ala arg val asp glu ala ala arg thr met his phe gly asp gly thr
                                        751/251
721/241
tee gaa eeg tte gat etg ett gee gtg gte eec eeg eac gtg eec tee gee geg eeg
ser glu pro phe asp leu leu ala val val pro pro his val pro ser ala ala ala arg
                                        811/271
781/261
tca gcg ggt ctc agc gaa tcc ggg tgg ata ccc gtg gac ccg cgc acc ctg tcc act agc
ser ala gly leu ser glu ser gly trp ile pro val asp pro arg thr leu ser thr ser
841/281
                                        871/291
gcc gac aac gtg tgg gcc atc ggc gat gcg acc gtg ctg acg ctg ccg aat ggc aaa ccg
ala asp asn val trp ala ile gly asp ala thr val leu thr leu pro asn gly lys pro
901/301
                                        931/311
ctg ccc aag gct gcc gtg ttc gcc gaa gcc cag gcc gca gtt gtc gcc cac ggc gtc gcc
leu pro lys ala ala val phe ala glu ala gln ala ala val val ala his gly val ala
                                        991/331
961/321
ege cat etc ggt tae gae gta get gag ege cae tte ace gge acg gge gee tge tae gte
arg his leu gly tyr asp val ala glu arg his phe thr gly thr gly ala cys tyr val
                                        1051/351
1021/341
qaq acc ggt gat cac cag gca gcc aag ggc gac ggc gat ttc ttc gct ccg tcg gcg ccc
glu thr gly asp his gln ala ala lys gly asp gly asp phe phe ala pro ser ala pro
                                         1111/371
1081/361
tcg gtg acg ctg tac ccg ccg tcg cgg gag ttt cac gag gag aag gtc gca caa gaa ctg
ser val thr leu tyr pro pro ser arg glu phe his glu glu lys val ala gln glu leu
1141/381
gcc tgg ctg acc cgc tgg aag acg tga
ala trp leu thr arg trp lys thr OPA
```

SEQ ID No.48D

FIGURE 48D



ORF according to Cole et al., 1998 (Nature 393: 537-544) and containing coding sequence Rv0331:

1/1									31/11	1								
tga aca	ccc	aca	ccq	acq	cqq	cqa	caa	tcg	cgg a	aaa	acc	ggt	ccq	cqq	gaa	tac	tac	aaa
OPA thr																		
61/21	•		•		-	-	-		91/31				•	-	•	4	2	5-1
cca tgg	gcc	gat	aat	agt	ttg	act	gac	tcg	gtc a	agt	cac	ccc	aag	acc	ttg	cgc	aag	act
pro trp	ala	asp	asn	ser	leu	thr	asp	ser	val s	ser	his	pro	lys	thr	leu	arg	lys	thr
121/41		-					_		151/5	51						-	-	
gcg gcg	gaa	tct	aat	att	cca	aag	ata	tat	gga a	act	cga	tgc	gaa	gga	atc	agg	ctc	atg
ala ala	glu	ser	asn	ile	pro	lys	ile	tyr	gly t	thr	arg	cys	glu	gly	ile	arg	leu	met
181/61									211/7	. —								
agc aag																		
ser lys	thr	val	leu	ile	leu	gly	ala	gly	val ç	gly	gly	leu	thr	thr	ala	asp	thr	leu
241/81									271/9									
cgt caa																		
arg gln	leu	leu	pro	pro	glu	asp	arg	ile			val	asp	arg	ser	phe	asp	gly	thr
301/101									331/1									
ctg ggc																		
leu gly	leu	ser	leu	leu	trp	val	leu	arg			arg	arg	pro	asp	asp	val	arg	val
361/121									391/1									
cgc ccc																		
arg pro	thr	ala	ala	ser	leu	pro	дтÀ	val			val	thr	ala	thr	val	ala	his	ıle
421/141									451/1								4.4	4
gac atc																		
asp ile	ата	ата	gru	vai	vaı	nis	unr	asp	511/1		vai	тте	дтй	CAL	asp	ата	reu	vaı
481/161	++-	~~+	~~~	~~~	a+ «	224	200	~~~			~~~	~~~	a+~	+ ~~	~~~	~~~	at a	~ ~ ~ ~
atc gca					_				_	-		-	_	-	-			-
ile ala 541/181	reu	дту	ala	ala	reu	asii	CIII	asp	571/1		PLO	gră	Teu	ser	asp	ата	reu	asp
gcc gac	ata	aaa	~~~	cac	ttc	tac	200	ata			aca	act	asa.	cta	cat	aca	224	ata
ala asp																		
601/201	Val	ala	9×3	9111	pne	CYL	CIII	100	631/2		ата	ата	gru	ıcu	ary	ara	T Y 2	Val
gag gcg	ctc	nan	cat	aac	caa	atc	act	ata	-		acc	aaa	ata	cca	ttc	aaa	tac	cca
qlu ala																		
661/221	200	9		9-1	~~9				691/2			5-1		1	F	1	-1-	F
gcc gca	cca	ttc	gaa	aca	aca	ttt	cta	atc			caa	ctc	aat	gac	cac	tac	acc	acc
ala ala																		
721/241	£	2	J						751/2		,			•	,	-		
gga acc	qta	cag	atc	gac	acg	ttc	acg	cct	gac	ccg	ctg	ccg	atg	ccc	gtt	gca	ggt	ccc
gly thr																		
781/261		-		•		-		-	811/			-		-				-
gag gtc	ggc	gag	gct	ttg	gtc	tcg	atg	ctc	aag (gat	cac	ggt	gtc	ggc	ttc	cat	cct	cgc
glu val																		
841/281		_							871/	291								
aag gcc	cta	gct	cgc	gtc	gat	gag	gcc	gca	agg a	acg	atg	cac	ttc	ggt	gac	ggc	acg	tcc
lys ala																		

SEQ ID No.48F

FIGURE 48F



931/311 901/301 gaa eeg tte gat etg ett gee gtg gte eee eeg eae gtg eee tee gee geg geg egg tea glu pro phe asp leu leu ala val val pro pro his val pro ser ala ala ala arg ser 991/331 961/321 geg ggt ctc age gaa tee ggg tgg ata eee gtg gae eeg ege ace etg tee aet age gee ala gly leu ser glu ser gly trp ile pro val asp pro arg thr leu ser thr ser ala 1051/351 1021/341 gac aac gtg tgg gcc atc ggc gat gcg acc gtg ctg acg ctg ccg aat ggc aaa ccg ctg asp asn val trp ala ile gly asp ala thr val leu thr leu pro asn gly lys pro leu 1111/371 1081/361 ccc aag gct gcc gtg ttc gcc gaa gcc cag gcc gca gtt gtc gcc cac ggc gtc gcc cgc pro lys ala ala val phe ala glu ala gln ala ala val val ala his gly val ala arg 1171/391 1141/381 cat ctc ggt tac gac gta gct gag cgc cac ttc acc ggc acg ggc gcc tgc tac gtc gag his leu gly tyr asp val ala glu arg his phe thr gly thr gly ala cys tyr val glu 1231/411 1201/401 acc ggt gat cac cag gca gcc aag ggc gac ggc gat ttc ttc gct ccg tcg gcg ccc tcg thr gly asp his gln ala ala lys gly asp gly asp phe phe ala pro ser ala pro ser 1291/431 1261/421 gtg acg ctg tac ccg ccg tcg cgg gag ttt cac gag gag aag gtc gca caa gaa ctg gcc val thr leu tyr pro pro ser arg glu phe his glu glu lys val ala gln glu leu ala 1321/441 tgg ctg acc cgc tgg aag acg tga trp leu thr arg trp lys thr OPA

SEO ID No.48F (continued)

FIGURE 48F (continued)



Fragment amplified by PCR based on the sequence similarities with a serine protease of the E.coli htrA family (creation of the BamHI site at the 5' end and of the SnaBI site at the 3' end) and subcloned into the vector pJVED:

31/11 1/1 cca tot aca ccg ctc aac ago cgg gcc aga cgc tgc cgg tcg gtg ctg ccg aga agg cgg pro ser thr pro leu asn ser arg ala arg cys arg ser val leu pro arg arg arg 91/31 tga tcc gtg gcg agt tgt tca tgt cgc ggc gca cca ccg ccg acc aac ggg tgc ttg cca OPA ser val ala ser cys ser cys arg gly ala pro pro pro thr asn gly cys leu pro 151/51 121/41 tee gte tga eea aeg gta gtt ege tge tga tet eea aaa gte tea age eea eeg aag eag ser val OPA pro thr val val arg cys OPA ser pro lys val ser ser pro pro lys gln 211/71 181/61 tca tga aca agc tgc gtt ggg tgc tat tga tcg tgg gtg gga tcg ggg tgg cgg tcg ccg ser OPA thr ser cys val gly cys tyr OPA ser trp val gly ser gly trp arg ser pro 271/91 241/81 cgg tgg ccg ggg gga tgg tca ccc ggg ccg ggc tga ggc cgg tgg gcc gcc tca ccg aag arg trp pro gly gly trp ser pro gly pro gly OPA gly arg trp ala ala ser pro lys 331/111 cgg ccg agc ggg tgg cgc gaa ccg acg acc tgc ggc cca tcc ccg tct tcg gca gcg acg arg pro ser gly trp arg glu pro thr thr cys gly pro ser pro ser ser ala ala thr 391/131 361/121 aat tgg cca ggc tga cag agg cat tca att taa tgc tgc ggg cgc tgg ccg agt cac ggg asn trp pro gly OPA gln arg his ser ile OCH cys cys gly arg trp pro ser his gly 451/151 aac ggc agg caa ggc tgg tta ccg acg ccg gac atg aat tgc gta ccc cgc taa cgt cgc asn gly arg gln gly trp leu pro thr pro asp met asn cys val pro arg OCH arg arg 511/171 481/161 tgc gca cca atg tcg aac tct tga tgg cct cga tgg ccc cgg ggg ctc cgc ggc tac cca cys ala pro met ser asn ser OPA trp pro arg trp pro arg gly leu arg gly tyr pro 571/191 age agg aga tgg teg ace tge gtg eeg atg tge tgg etc aaa teg agg aat tgt eea cae ser arg arg trp ser thr cys val pro met cys trp leu lys ser arg asn cys pro his 631/211 601/201 tgg tag gcg att tgg tgg acc tgt ccc gag gcg acg ccg gag aag tgg tgc acg agc cgg trp AMB ala ile trp trp thr cys pro glu ala thr pro glu lys trp cys thr ser arg 691/231 661/221 teg aca tgg ctg acg teg teg acc gca gcc tgg age ggg tca gge gge gca acg ata ser thr trp leu thr ser ser thr ala ala trp ser gly ser gly gly gly ala thr ile 751/251 721/241 tcc ttt tcg acg tcg agg tga ttg ggt ggc agg ttt atg gcg ata ccg ctg gat tgt cgc ser phe ser thr ser arg OPA leu gly gly arg phe met ala ile pro leu asp cys arg 811/271 781/261 gga tgg cgc tta acc tga tgg aca acg ccg cga agt gga gcc cgc cgg gcg gcc acg tgg gly trp arg leu thr OPA trp thr thr pro arg ser gly ala arg arg ala ala thr trp 871/291 841/281 gtg tca ggc tga gcc agc tcg acg cgt cgc acg ctg agc tgg tgt tcc ccg acc gcg gcc val ser gly OPA ala ser ser thr arg arg thr leu ser trp trp phe pro thr ala ala

SEO ID No.49A

FIGURE 49A

HER A CONSESSION CERTIFIC CERTIFIED OF

								001									
901/301								931/			~~+	+++	200	aat	aaa	cat	caa
cgg gca ttc	ccg	tgc	agg	agc	gcc	gtc	tgg	tgt	100	aac	ggc	nha	thr	alu	ara	his	ara
arg ala phe	pro	cys	arg	ser	aıa	vaı	trp	991/	1eu '331	asii	gry	pne	CIII	gry	arg	1113	arg
961/321 cac ggg cgt	tac	caa	att	caa	acc	t.ca	aat			tca	tca	aac	agg	tgg	tgc	tca	acc
his gly arg	CVS	ara	val	arg	ala	ser	alv	trp	arg	ser	ser	asn	arg	trp	cys	ser	thr
1021/341	Cyb	urg	•	429			3-1	1051	L/35	1			-	-	_		
aca aca aat	tac	tac	σca	tcq	aaq	aca	ccg	acc	cag	gcg	gcc	agc	ccc	ctg	gaa	cgt	cga
thr ala asp	cys	cys	ala	ser	lys	thr	pro	thr	gln	ala	ala	ser	pro	leu	glu	arg	arg
1081/361								1111	L/37	<u>l</u>							
ttt acd tdc	tgc	tcc	ccg	gcc	gtc	gga	tgc	cga	ttc	cgc	agc	ttc	ccg	gtg	cga	cgg	ctg
phe thr cys	cys	ser	pro	ala	val	gly	cys	arg	phe	arg	ser	phe	pro	val	arg	arg	leu
1141/381								1171	1/39	1							
gcg ctc gga	gca	cgg	aca	tcg	aga	act	ctc	ggg	gtt	cgg	cga	acg	tta	tct	cag	tgg	aat
ala leu gly	ala	arg	thr	ser	arg	thr	leu	gly	val	arg	arg	thr	leu	ser	gın	trp	asn
1201/401									1/41						+	~~~	-
ctc agt cca	cgc	gcg	caa	cct	agt	tgt	gca	gtt	act	gtt	gaa	agc	cac	acc +hr	hic	313	agt
leu ser pro	arg	ala	gln	pro	ser	cys	ala	val	tnr 1/43	vaı 1	gru	ser	HITS	CILL	1117.2	ата	261
1261/421											C 3 C	~ a a	asa	caa	cct	agc	gac
cca cgc atg	gcc	aag	ttg	gcc	cga	gta	gra	ggc	lan	y ca val	aln	alii	alu	aln	pro	ser	asp
	aıa	туѕ	Leu	ата	arg	vaı	Val	9±y	$\frac{1}{45}$	1	9	gra	9-4	9	F		F
1321/441 atg acg aat		999	~~~	+ > +	taa	cca	cca	בכם	cad	cao	cca	gga	acc	cca	aat	tat	qct
met thr asn	hic	nro	ara	tur	ser	pro	pro	pro	aln	gln	pro	qly	thr	pro	gly	tyr	āla
1381/461	1113	PLO	arg	012		P	F	141	1/47	1	•	-		_		-	
מפת תתת כפת	cag	caa	acq	tac	agc	cag	cag	ttc	gac	: tgg	cgt	tac	cca	ccg	tcc	ccg	CCC
gln gly gln	aln	gln	thr	tyr	ser	gln	gln	phe	asp	trp	arg	tyr	pro	pro	ser	pro	pro
1441/481								147	1/49	1							
cca caa cca	acc	cag	tac	cgt	caa	ccc	tac	gag	gcg	, ttg	ggt	ggt	acc	cgg	ccg	ggt	ctg
pro gln pro	thr	gln	tyr	arg	gln	pro	tyr	glu	ala	ı leu	gly	gly	thr	arg	pro	āΤλ	leu
1501/501								153	1/51	.1							
ata cct ggc	gtg	att	ccg	acc	atg	acg	ccc	: cct	cct	ggg	atg	gtt	cgc	caa	. cgc	CCT	. cgt
ile pro gly	val	ile	pro	thr	met	thr	pro	pro	pro	o gră	met	vaı	arg	gin	arg	pro	arg
1561/521									1/53		. a+a	too	~~~		ato	. ממכ	. aac
gca ggc atg	ttg	gcc	atc	ggc	gcg	gtg	acq	, ala	ge	y gre	y gry	ser	gcc ala	alt.	, ale	alv	ggo alv
ala gly met	leu	ala	ıle	g T Y	ата	Val	. CIII	165	$\frac{1}{5}$	1 Val	. vai	301	ити	9-1		9-1	9-1
1621/541 gcg gcc gca			~+~		++-						r aac	ccc	ago	aac	aac	cca	qtq
gcg gcc gca ala ala ala	tcc	101	guo	999 710	nhe	act	arc	, gco	pro	o ala	a alv	pro	ser	gly	gly	pro	val
	Ser	reu	val	. gry	piic	. ubi		171	1/5	71	- 5-1	•		-		_	
1681/561 gct gcc ago		י מכמ	cca	ago	ato	ccc	gea				g ccg	ccg	ggg	, tc	gto	gaa	a cag
ala ala ser	ala - ala	ala	pro	ser	ile	pro	ála	a ála	aas	n met	pro	pro	gly	, sei	r val	. glu	ı gln
17/1/581								177	71/5	91							
~+~ ~~~ ~~	aac	r ata	gto	ccc	agt	gto	gto	c ato	g tt	g gaa	a acc	gat	cto	g ggo	c cgc	caq	g tcg
val ala ala	lys	val	val	pro	ser	val	l vai	l met	t le	u gii	u thi	asp) lei	ıgl	yarq	g gli	n ser
1001/601								183	31/6	11							
~~~ ~~~ ~~	tcc	ggc	ato	att	cto	g tct	t gc	c gaç	g gg	g cto	g ato	tto	g acc	c aa	c aad	c ca	c gtg
glu glu gly	y sei	gly	, il∈	e ile	e leu	ı se	r ala	a glı	ı g⊥	у Ге	u ile	e let	ı thi	r ası	n ası	n ni:	s var
1861/621						_			91/6		~ ~~	*		7 20	n arti	a	
atc gcg gcg	g gç	gcc	aaq	g cct	ccc	cto	g gg	c agi	CC	g cc	y ccg	y dad	a acc	y ac	y yc	^ I	
ile ala ala	a ala	a ala	lys	s pro	pro	) Lei	u gi	y se:	r pr	o pr	Opro	, TA:	o CIII	L CII.	. va.	-	

SEQ ID No.49A (continued 1)

FIGURE 49A (continued 1)

31/11 cat cta cac cgc tca aca gcc ggg cca gac gct gcc ggt cgg tgc tgc cga gaa ggc ggt his leu his arg ser thr ala gly pro asp ala ala gly arg cys cys arg glu gly gly 91/31 gat ccg tgg cga gtt gtt cat gtc gcg gcg cac cac cgc cga cca acg ggt gct tgc cat asp pro trp arg val val his val ala ala his his arg arg pro thr gly ala cys his 121/41 151/51 ccg tct gac caa cgg tag ttc gct gct gat ctc caa aag tct caa gcc cac cga agc agt pro ser asp gln arg AMB phe ala ala asp leu gln lys ser gln ala his arg ser ser 211/71 cat gaa caa gct gcg ttg ggt gct att gat cgt ggg tgg gat cgg ggt ggc ggt cgc cgc his glu gln ala ala leu gly ala ile asp arg gly trp asp arg gly gly gly arg arg 271/91 241/81 ggt ggc cgg ggg gat ggt cac ccg ggc cgg gct gag gcc ggt ggg ccg cct cac cga agc gly gly arg gly asp gly his pro gly arg ala glu ala gly gly pro pro his arg ser 331/111 301/101 ggc cga gcg ggt ggc gcg aac cga cga cct gcg gcc cat ccc cgt ctt cgg cag cga cga gly arg ala gly gly ala asn arg arg pro ala ala his pro arg leu arg gln arg arg 391/131 361/121 att ggc cag gct gac aga ggc att caa ttt aat gct gcg ggc gct ggc cga gtc acg gga ile gly gln ala asp arg gly ile gln phe asn ala ala gly ala gly arg val thr gly 451/151 acg gca ggc aag gct ggt tac cga cgc cgg aca tga att gcg tac ccc gct aac gtc gct thr ala gly lys ala gly tyr arg arg thr OPA ile ala tyr pro ala asn val ala 511/171 481/161 gcg cac caa tgt cga act ctt gat ggc ctc gat ggc ccc ggg ggc tcc gcg gct acc caa ala his gln cys arg thr leu asp gly leu asp gly pro gly gly ser ala ala thr gln 571/191 gca gga gat ggt cga cct gcg tgc cga tgt gct ggc tca aat cga gga att gtc cac act ala gly asp gly arg pro ala cys arg cys ala gly ser asn arg gly ile val his thr 631/211 601/201 ggt agg cga ttt ggt gga cct gtc ccg agg cga cgc cgg aga agt ggt gca cga gcc ggt gly arg arg phe gly gly pro val pro arg arg arg arg ser gly ala arg ala gly 691/231 cga cat ggc tga cgt cgt cga ccg cag cct gga gcg ggt cag gcg gcg caa cga tat arg his gly OPA arg arg arg pro gln pro gly ala gly gln ala ala ala gln arg tyr 751/251 721/241 cet ttt cga cgt cga ggt gat tgg gtg gca ggt tta tgg cga tac cgc tgg att gtc gcg pro phe arg arg gly asp trp val ala gly leu trp arg tyr arg trp ile val ala 811/271 gat ggc gct taa cct gat gga caa cgc cgc gaa gtg gag ccc gcc ggg cgg cca cgt ggg asp gly ala OCH pro asp gly gln arg arg glu val glu pro ala gly arg pro arg gly 871/291 841/281 tgt cag gct gag cca gct cga cgc gtc gca cgc tga gct ggt ggt ttc cga ccg cgc ccc cys gln ala glu pro ala arg arg val ala arg OPA ala gly gly phe arg pro arg pro 931/311 901/301 ggg cat tee egt gea ggg eeg tet ggt gtt tga acg gtt tta eeg gte gge ate gge gly his ser arg ala gly ala pro ser gly val OPA thr val leu pro val gly ile gly 991/331 961/321 acg ggc gtt gcc ggg ttc ggg cct cgg gtt ggc gat cgt caa aca ggt ggt gct caa cca thr gly val ala gly phe gly pro arg val gly asp arg gln thr gly gly ala gln pro 1051/351 1021/341 cgg cgg att gct gcg cat cga aga cac cga ccc agg cgg cca gcc ccc tgg aac gtc gat arg arg ile ala ala his arg arg his arg pro arg arg pro ala pro trp asn val asp

SEO ID No.49B

FIGURE 49B

1081/361	1111/371
tta cgt gct gct ccc cgg ccg tcg gat gcc	gat tee gea get tee egg tge gae gge tgg
leu arg ala ala pro arg pro ser asp ala	asp ser ala ala ser arg cys asp gly trp
1141/381	1171/391
cgc tcg gag cac gga cat cga gaa ctc tcg	ggg ttc ggc gaa cgt tat ctc agt gga atc
arg ser glu his gly his arg glu leu ser	gly phe gly glu arg tyr leu ser gly ile
1201/401	1231/411
tca gtc cac gcg cgc aac cta gtt gtg cag	tta ctg ttg aaa gcc aca ccc atg cca gtc
	leu leu lys ala thr pro met pro val
1261/421	1291/431
cac gca tgg cca agt tgg ccc gag tag tgg	gcc tag tac agg aag agc aac cta gcg aca
	ala AMB tyr arg lys ser asn leu ala thr
1321/441	1351/451
tga cga atc acc cac ggt att cgc cac cgc	cgc agc agc cgg gaa ccc cag gtt atg ctc
. <del>-</del>	arg ser ser arg glu pro gln val met leu 1411/471
1381/461	tog act ggo gtt acc cac cgt ccc cgc ccc
agg ggc agc agc aad cgc aca gcc agc agc	ser thr gly val thr his arg pro arg pro
arg gry ser ser rys arg thr ara ser ser 1441/481	1471/491
cad add caa eed adt add dto aac eet aco	agg cgt tgg gtg gta ccc ggc cgg gtc tga
arg ser old noo ser thr val asm pro thr	arg arg trp val val pro gly arg val OPA
1501/501	1531/511
tac ctg gcg tga ttc cga cca tga cgc ccc	ctc ctg gga tgg ttc gcc aac gcc ctc gtg
tyr leu ala OPA phe arg pro OPA arg pro	leu leu gly trp phe ala asn ala leu val
1561/521	1591/531
	tag egg tgg tgt eeg eeg gea teg geg geg
gln ala cys trp pro ser ala arg OPA arg	AMB arg trp cys pro pro ala ser ala ala
1621/541	1651/551
cgg ccg cat ccc tgg tcg ggt tca acc ggg	cac ccg ccg gcc cca gcg gcg gcc cag tgg
arg pro his pro trp ser gly ser thr gly	his pro pro ala pro ala ala ala gln trp
1681/561	1711/571
ctg cca gcg cgg cgc caa gca tcc ccg cag	caa aca tgc cgc cgg ggt cgg tcg aac agg
leu pro ala arg arg gln ala ser pro glr	gln thr cys arg arg gly arg ser asn arg
1741/581	1771/591
tgg cgg cca agg tgg tgc cca gtg tcg tca	tgt tgg aaa ccg atc tgg gcc gcc agt cgg
·	cys trp lys pro ile trp ala ala ser arg
1801/601	1831/611
agg agg gct ccg gca tca ttc tgt ctg ccg	g agg ggc tga tct tga cca aca acc acg tga
	arg gly OPA ser OPA pro thr thr thr OPA 1891/631
1861/621	
tcg cgg cgg ccg cca agc ctc ccc tgg gca ser arg arg pro pro ser leu pro trp ala	a val ard ard ard lys ard ard
ser and and bro bro ser red bro crb are	, tar ard ard ard alo ard ard

SEQ ID No.49B (continued 1)

FIGURE 49B (continued 1)



1/1				31/11						
atc tac acc gct	caa cag	ccg ggc	cag acg	ctg ccg	gtc ggt	gct g	gcc gag	aag	gcg	gtg
ile tyr thr ala 61/21	gln gln	pro gly	gln thr	leu pro 91/31	val gly	ala a	ala glu	lys	ala	val
atc cgt ggc gag										
ile arg gly glu 121/41	leu phe	met ser	arg arg	thr thr a 151/51	ala asp	gln a	arg val	leu	ala	ile
cgt ctg acc aac	ggt agt	tcg ctg	ctg atc	tcc aaa a	agt ctc	aag d	ccc acc	gaa	gca	gtc
arg leu thr asr 181/61				211/71						
atg aac aag ctg										
met asn lys leu 241/81				271/91						
gtg gcc ggg ggg							~		_	, ,
val ala gly gly 301/101				331/111					_	
gcc gag cgg gtg										
ala glu arg val 361/121				391/131						
ttg gcc agg ctg										
leu ala arg leu 421/141		_		451/151					_	_
cgg cag gca agg										
arg gln ala arg 481/161				511/171						
cgc acc aat gto										
arg thr asn val 541/181				571/191						
cag gag atg gto										
gln glu met vai 601/201				631/211						
gta ggc gat tto										
val gly asp let 661/221				691/231						
gac atg gct gad										
asp met ala asp 721/241				751/251						
ctt ttc gac gt										
leu phe asp val 781/261				811/271						
atg gcg ctt aa										
met ala leu asi 841/281		_		871/291						
gtc agg ctg ag	c cag ctc	gac gcg	tcg cac	gct gag	ctg gtg	gtt	tcc gac	cgc	ggc	ccg
val arg leu se 901/301				931/311						
ggc att ccc gt	g cag gag	cgc cgt	ctg gtg	ttt gaa	cgg ttt	tac	cgg tcg	gca	tcg	gca
gly ile pro va 961/321				991/331						
cgg gcg ttg cc										
arg ala leu pr	o gry ser	gry reu	gry reu	ara ire	var rys	grn	var Val	reu	asn	nıs

SEQ ID No.49C

FIGURE 49C



								1051	1/25								
1021/341							~~~		1/35		cad	CCC	cct	aaa	acq	tica	att
ggc gga ttg	ctg	cgc	atc	gaa	gac	acc +h-	gac	nra	99C	al v	aln	nro	nro	alv	thr	ser	ile
gly gly leu	leu	arg	ше	gru	asp	CIII	asp	1111	9±3 L/37:	g±y I	9-11	Pro	P	9-1			
1081/361 tac gtg ctg			~~~	aat	~~~	ata	cca				ctt	ccc	aat	aca	acq	act	aac
tac gtg ctg tyr val leu	CTC	ccc	ggc	ara	ara	met	nro	ile	pro	aln	leu	pro	alv	ala	thr	āla	gly
	reu	pro	g r y	arg	arg	mee	PLU	117:	1/39	1		•					
1141/381 gct cgg agc	a C (1	aac	atc	σασ	aac	tct	caa	aat	tcg	gcg	aac	gtt	atc	tca	gtg	gaa	tct
ala arg ser	thr	asp	ile	alu	asn	ser	arg	gly	ser	ala	asn	val	ile	ser	val	glu	ser
1201/401								123	1/41	1							
and tag aca	cac	qca	acc	tag	ttg	tgc	agt	tac	tgt	tga	aag	cca	cac	cca	tgc	cag	tcc
gln ser thr	arq	ala	thr	AMB	leu	cys	ser	tyr	cys	OPA	lys	pro	his	pro	cys	gln	ser
1261/421								129	1/43	1							
acg cat ggc	caa	gtt	ggc	ccg	agt	agt	ggg	cct	agt	aca	gga	aga	gca	acc	tag	cga	cat
thr his gly	gln	val	gly	pro	ser	ser	gly	pro	ser	thr	gly	arg	ala	thr	AMB	arg	nıs
1221/441								135	1/40	1							
gac gaa tca	CCC	acg	gta	ttc	gcc	acc	gcc	gca	gca	gcc	ggg	aac	CCC	agg	lou	cyc	cer
asp glu ser	pro	thr	val	phe	ala	thr	ala	ala	ala	ala 1	дтА	asn	bro	arg	Teu.	Cys	261
1381/461									1/47		++=	acc	200	ata	ccc	acc	CCC
ggg gca gca	gca	aac	gta	cag	cca	gca	gtt	cga	1019	212	len	nro	thr	val	pro	ala	pro
gly ala ala	ala	asn	val	gın	pro	ата	val	117	1/49	. ата 1	reu	pro	CIII	Vul	PLU	<b>u_</b> u	PI
1441/481 gca gcc aac				<b>.</b>		at a	aaa				taa	tac	cca	acc	aaa	tct	gat
gca gcc aac ala ala asn	cca	gta	ccg	cor	thr	len	ard	ggc	val	alv	tro	tvr	pro	ala	qly	ser	asp
	pro	Val	pro	261	CIII	104	u_9	153	1/51	1		-	•		-		_
1501/501 acc tgg cgt	ant.	tcc	aac	cat	gac	acc	ccc				ggt	tcg	cca	acg	ccc	tcg	tgc
thr trp arg	. yac	ser	asp	his	asp	ala	pro	ser	tr	asp	gly	ser	pro	thr	pro	ser	cys
1561/501								159	)1/5:	31							
	aac	cat	caa	cgc	ggt	gac	: gat	ago	ggt	: ggt	gtc	cgc	cgg	cat	cgg	cgg	cgc
arg his val	. alv	his	arg	arg	gly	asp	asp	ser	gly	gly	val	arg	arg	his	arg	arg	arg
1601/5/1								165	) 1 / D	) I							
ggc cgc ato	cct	ggt	cgg	gtt	caa	. ccg	ggq	acc	cg	cgg	ccc	cag	cad	l cad	ccc	agt	ggc
gly arg ile	e pro	gly	arg	val	. glr	pro	gly	tui	rar	garo	g pro	gin	arç	garo	pro	ser	g gry
1601/561								1/2	LT/ 2	<i>1</i> 1							
tgc cag cgc	ggc	gcc	aag	cat	ccc	cgc	ago	aaa	a cai	z gco	gcc	ggg	gto	ggt	cga	. aca	ggt - alv
cys gln arg	g gly	, ala	lys	his	pro	arç	g sei	r TA	s hi:	5 ala	a ala	ı g⊥y	/ Val	91)	arg	CIII	. gry
1741/581								1/	71/5	91 		. ++	- ~~		1 CC8	ato	r dda
ggc ggc caa	a ggt	: ggt	gco	cac	g tgt	. cgt	ca:	c gc	t gg	a aau	o erc	r sei	- 999	nro	pro	val	alv
gly gly gli	n gly	, gly	ala	ı gır	ı cys	arç	i ur:	o Val. 1Ω'	1 91 31/6	y ası 11	. arc	,	- y-:	, ,,,			- <u>5</u> -1
1801/601 gga ggg ct				- +~+	- ~+	- +~	~ ~~				t ett	ga c	c caa	a caa	a cca	a cat	gat
gga ggg cto gly gly le	c cgg	g cat	, cat		- yu	- cy	s are	יפפי	y yc	a ası	o lei	ası	o ql	n qli	n pro	aro	asp
	u arç	1112	) !IT?	9 G T	_ va.	_ ~у.		18	91/6	31				,	-	•	_
1861/621 cgc ggc gg	c ca	~ Ca=	א מכי	a taa	a def	t aa	g ca				c gaa	a aac	c ga	c gg	t a		
arg gly gl	u are	r alr	. yo\ nala	sei	r pr	o ali	v al	n se	r al	a al	a glı	ı ası	n as	p gl	У		
arg gry gr	y ard	9 911			- 2-,	- 9-	, , , -				-		•				

SEQ ID No.49C (continued 1)

FIGURE 49C (continued 1)



Coding sequence Rv0983 predicted by Cole et al., 1998 (Nature 393:537-544) and containing seq60A:

1/1										31/1	1								
atg g	cc a	aaq	ttg	gcc	cga	gta	gtg	ggc	cta	gta	cag	gaa	gag	caa	cct	agc	gac	atg	acg
Met a	la :	lys	leu	ala	arg	val	val	gly	leu	val	gln	glu	glu	gln	pro	ser	asp	met	thr
61/21										91/3	1								
aat c	ac ·	cca	cgg	tat	tcg	cca	ccg	ccg	cag	cag	ccg	gga	acc	cca	ggt	tat	gct	cag	ggg
asn h	is ;	pro	arg	tyr	ser	pro	pro	pro	gln	gln	pro	gly	thr	pro	gly	tyr	ala	gln	gly
121/4	1	_								151/									
cag c	ag	caa	acg	tac	agc	cag	cag	ttc	gac	tgg	cgt	tac	cca	ccg	tcc	ccg	ccc	ccg	cag
gln g	ln	gln	thr	tyr	ser	gln	gln	phe	asp	trp	arg	tyr	pro	pro	ser	pro	pro	pro	gln
181/6	1									211/									
cca a	cc	cag	tac	cgt	caa	CCC	tac	gag	gcg	ttg	ggt	ggt	acc	cgg	ccg	ggt	ctg	ata	cct
pro t	hr	gln	tyr	arg	gln	pro	tyr	glu	ala	leu	gly	gly	thr	arg	pro	gly	leu	ile	pro
241/8										271/									
ggc g	tg	att	ccg	acc	atg	acg	ccc	cct	cct	ggg	atg	gtt	cgc	caa	cgc	cct	cgt	gca	aac
gly v	al	ile	pro	thr	met	thr	pro	pro	pro			val	arg	gīn	arg	pro	arg	ala	дтА
301/1	01									331/									
atg t	tg	gcc	atc	ggc	gcg	gtg	acg	ata	gcg	gtg	gtg	tcc	gcc	ggc	atc	ggc	ggc	gcg	gcc
met 1	eu	ala	ile	gly	ala	val	thr	ile	ala			ser	ala	āтĀ	ııe	дтА	дтĀ	aıa	ala
361/1	21									391/							4		
gca t	CC	ctg	gtc	ggg	ttc	aac	cgg	gca	ccc	gcc	ggc	CCC	agc	ggc	ggc	cca	gcg	gct	gcc
ala s		leu	val	gly	phe	asn	arg	ala	pro			pro	ser	дтĀ	дту	bro	vai	ата	ala
421/1										451/			~~~	+ ~ ~	~+ ~	~~~	a24	~+~	~~~
agc g	lcd	gcg	cca	agc	atc	ccc	gca	gca	aac	acg	ccg	ccg	999	cor	g	gaa	aln	ycy	9C9
ser a		ala	pro	ser	ııe	pro	aıa	aıa	asn	met 511/		bro	gry	Ser	vaı	gru	gin	vaı	ara
481/1 gcc a	.61	4		~~~	a ##	ata	a+ a	ata	++~			cat	cta	aac	cac	cad	tca	gag	gag
gcc a ala l	lag	gtg	gug	200	agt	900	3721	mat	lau	gaa	thr	asn	leu	alv	arg	aln	ser	alu	alu
		vaı	vai	рго	ser	vai	Val	mec	Teu		/191	азр	104	9-1	arg	9		9-4	9
541/1 ggc t	81		-+-	~ <del>+ +</del>	ata	+ ~+	~~~	asa	aaa			tta	acc	aac	aac	cac	ata	atc	aca
ggc t	cc	ggc	ila	ila	lau	car	900 ala	gay	999	len	ile	len	thr	asn	asn	his	val	ile	ala
		gry	TTG	TIC	Ieu	261	ara	gra	9 T Y		/211	1.0u		u					
601/2 gcg g	701	~~~	220	cct	ccc	cta	ממכ	agt	cca			aaa	acq	aca	gta	acc	ttc	tct	gac
ala a	112	313	luc	nro	nro	leu	alv	ser	nro	nro	pro	lvs	thr	thr	val	thr	phe	ser	asp
661/2		ата	TÀS	pro	Pro	ıcu	9+1	501	PIO	691	/231	-1-							-
ggg c	227	200	aca	ccc	ttc	acq	ata	ata	aaa				acc	agt	gat	atc	qcc	qtc	gtc
gly a	-yy	thr	ala	pro	phe	thr	val	val	alv	ala	asp	pro	thr	ser	asp	ile	āla	val	val
721/2		CIII	ulu	PIO	P0	0112			5-1	751	/251	•			•				
cat c	rt.t.	car	aac	atc	tcc	aaa	ctc	acc	ccq	atc	tcc	ctg	ggt	tcc	tcc	tcg	gac	ctg	agg
arg v	val	aln	alv	val	ser	qlv	leu	thr	pro	ile	ser	leu	gĺy	ser	ser	ser	asp	leu	arg
781/2	261									811	/271								
atc o	aat	caq	cca	qtq	ctq	gcg	atc	ggg	tcg	ccg	ctc	ggt	ttg	gag	ggc	acc	gtg	acc	acg
val	gly	gln	pro	val	leu	ala	ile	gly	ser	pro	leu	gly	leu	glu	gly	thr	val	thr	thr

SEQ ID No.49D

FIGURE 49D

841/281								871/									
ggg atc gt	c ago	gct	ctc	aac	cgt	cca	gtg	tcg	acg	acc	ggc	gag	gcc	ggc	aac	cag	aac
gly ile va	ıl ser	ala	leu	asn	arg	pro	val			thr	gly	glu	ala	gly	asn	gln	asn
901/301								931/									
acc gtg ct	g gad	gcc	att	cag	acc	gac	gcc	gcg	atc	aac	ccc	ggt	aac	tcc	ggg	ggc	gcg
thr val le	eu asp	ala	ile	gln	thr	asp	ala			asn	pro	gly	asn	ser	gly	gly	ala
961/321								991/	-								
ctg gtg aa	ic atq	g aac	gct	caa	ctc	gtc	gga	gtc	aac	tcg	gcc	att	gcc	acg	ctg	ggc	gcg
leu val as	n met	asn	ala	gln	leu	val	gly				ala	ile	ala	thr	leu	gly	ala
1021/341 1051/351  gac tca gcc gat gcg cag agc ggc tcg atc ggt ctc ggt ttt gcg att cca gtc gac cag																	
gac tca go	c gat	gcg	cag	agc	ggc	tcg	atc	ggt	ctc	ggt	ttt	gcg	att	cca	gtc	gac	cag
asp ser al	a ası	ala	gln	ser	gly	ser	ile				phe	ala	ıle	pro	val	asp	gln
1081/361									1/37:								
gcc aag co	gc at	gcc	gac	gag	ttg	atc	agc	acc	ggc	aag	gcg	tca	cat	gcc	tcc	ctg	ggt
ala lys ar	g ile	e ala	asp	glu	leu	ile	ser				ala	ser	nıs	ala	ser	leu	дтА
1141/381									1/39:	_							
gtg cag gt	g ac	c aat	gac	aaa	gac	acc	ctg	gāc	gcc	aag	atc	gtc	gaa	gta	gtg	gcc	ggt
val gln va	al th	r asn	asp	lys	asp	thr	leu				11e	val	gru	val	vaı	ala	grā
1201/401									1/41		,						
ggt gct gc	cc gc	g aac	gct	gga	gtg	ccg	aag	ggc	gtc	gtt	gtc	acc	aag	gtc	gac	gac	cgc
gly ala al	la al	a asn	ala	дтЛ	val	pro	Tys				vaı	thr	туs	vaı	asp	asp	arg
1261/421									1/43		4					~~~	
ccg atc as	ac ag	c gcg	gac	gcg	ttg	gtt	gcc	gcc	gtg	cgg	tcc	aaa	gcg	ccg	ggc	gcc	acg
pro ile as	sn se	r ala	asp	ala	leu	vaı	ата		vai 1/45		ser	rys	ата	pro	gry	ala	CHL
1321/441									-		2.52	~+~	<b>a</b> aa	at a	200	ctc	aac
gtg gcg ct	ta ac	c ttt	cag	gat	CCC	ccg	ggc	ggt	age	cgc	aca +h-	gra	~laa	ycc 1	thr	100	ggc ~l··
val ala le	eu th	r phe	gın	asp	pro	ser	дтА	дту	ser	arg	UIII	Val	gin	val	CIII	reu	gry
1381/461																	
aag gcg ga	-																
lys ala gi	ıu gi	n OPA															

SEQ ID No.49D (continued 1)

FIGURE 49D (continued 1)



ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv0983

1/	1									31/1									
t a	a acc	agc	tcg	acg	cgt	cgc	acg	ctg	agc	tgg	tgg	ttt	ccg	acc	gcg	gcc	cgg	gca	ttc
OP	A ala	ser	ser	thr	arg	arg	thr	leu	ser	trp	trp	phe	pro	thr	ala	ala	arg	ala	phe
61	/21									91/3			~~+	~~~	ast	aaa	C 3 C	aaa	cat
CC	g tgc	agg	agc	gcc	gtc	tgg	tgt	ttg	aac	ggt	nhe	thr	ggu	ara	his	ara	his	alv	ara
-	_	arg	ser	ala	vaı	trp	cys	reu	asıı	151/	51	CIII	gry	arg	1115	u_g		9-1	urg
to	1/41 c cgg	att	caa	acc	t.ca	aat	taa	cqa	tcg	tca	aac	agg	tgg	tgc	tca	acc	acg	gcg	gat
C.	s arg	val	arq	ala	ser	gly	trp	arg	ser	ser	asn	arg	trp	cys	ser	thr	thr	ala	asp
1.5	1/61									211/	71								
to	c tgc	gca	tcg	aag	aca	ccg	acc	cag	gcg	gcc	agc	CCC	ctg	gaa	cgt	cga	ttt	acg	tgc
	s cys	ala	ser	lys	thr	pro	thr	gln	ala	a1a 271/	ser	pro	Ieu	gru	arg	arg	pne	CHL	Cys
24	11/81 jc tcc			~+~	~~~	taa	cas	ttc	cac			cca	ata	сда	caa	cta	aca	ctc	gga
to	s ser	ccg	gcc	guc	gga	CVS	ara	phe	arg	ser	phe	pro	val	arq	arg	leu	ala	leu	gly
3 (	11/101									331/	111								
~	ים כמת	aca	tca	aga	act	ctc	ggg	gtt	cgg	cga	acg	tta	tct	cag	tgg	aat	ctc	agt	cca
a.	La arg	thr	ser	arg	thr	leu	gly	val	arg	arg	thr	leu	ser	gln	trp	asn	leu	ser	pro
36	51/121									391/	131						_		
C	gc gcg	caa	cct	agt	tgt	gca	gtt	act	gtt	gaa	agc	cac	acc	cat	gcc	agt	cca	cgc	atg
	rg ala		pro	ser	cys	ala	val	thr	val	g1u 451/	ser /151	nıs	thr	nis	ala	ser	pro	ary	mec
4:	21/141 cc aag		~~~		a+ a	ata	aac	cta	αta			gag	caa	cct	agc	σac	atq	acq	aat
g	cc aag la lys	leu	gcc	ara	val	val	alv	leu	val	gln	glu	glu	gln	pro	ser	asp	met	thr	asn
4	81/161									511,	/1/1								
~		caa	tat	tcg	cca	ccg	ccg	cag	cag	ccg	gga	acc	cca	ggt	tat	gct	cag	ggg	cag
h	is pro	arg	tyr	ser	pro	pro	pro	gln	gln	pro	gra	thr	pro	gly	tyr	ala	gln	дтй	gin
5	41/181								4		/191	999	222	taa	ccc	ccc	cca	cad	CCa
С	ag caa	acg	tac	ago	cag	cag	ttc	gac	tgg	cgt	tac	nro	nro	ser	nro	nro	pro	aln	pro
	ln gln		tyr	ser	gın	gın	pne	asp	сгр	631	/211	pro	PLO	501	PLU	Pro	PIO	9	F
_	01/201 cc caç	+ + > 0	cat	caa	CCC	tac	σασ	aca	tta	aat	aat	acc	: cgg	ccg	ggt	ctg	ata	cct	ggc
a t	hr glr	tvr	ard	. aln	pro	tyr	glu	ala	leu	gly	gĺy	thr	arg	pro	gly	leu	ile	pro	gly
6	61/221									691	/231								
g	tg att	ccg	acc	atg	acg	ccc	cct	cct	ggg	atg	gtt	cgc	caa	cgc	cct	cgt	gca	ggc	atg
v	al ile	pro	thr	met	thr	pro	pro	pro	gly	met	val	arg	1 gin	arg	pro	arg	aıa	дту	mec
7	21/241	-									/251			ato	aac	. ממכ	aca	acc	gca
t	tg gco eu ala	ato	ggc	gcg	gtg	acg	ata	gcg	yes	y y cy val	ser	ala ala	alv	ile	alv	gg.	ala	ala	ala
	eu ala 81/263		* <b>Δ</b> τ?	, arg	ı val	LIII				811	/271		- y <b>-</b> 1		ړ – ر	J 1			
+	ac ata	r ata	a aa	ı tta	aac	cac	qca	ccc	gcc	qqq	ccc	ago	ggc	ggc	cca	gtg	gct	gco	agc
	er le	, you	r ali	phe	asr	arç	ala	pro	āla	ı gly	pro	sei	c gly	gly	pro	val	. ala	ala	ser
	41/20	1								871	/291	_							
ç	cg gc	g cca	agg	c ato	ccc	gca	gca	aac	ato	g ccg	CCG	ggg	g tcc	gto	: gaa	cag	gto	gco داد	gcc
ā	la ala	a pro	sei	r ile	pro	ala	a ala	asr	n met	pro	pro	dT.	y sei	. val	. gr	4 911	, val		

SEQ ID No.49F

FIGURE 49F

901/301								931/									
sea ata ata	CCC	agt	gtc	gtc	atg	ttg	gaa	acc	gat	ctg	ggc	cgc	cag	tcg	gag	gag	ggc
lys val val	pro	ser	val	val	met	leu	glu	thr	asp	leu	gly	arg	gln	ser	glu	glu	άīλ
961/321 tcc ggc atc					~~~	~~~		991/		200	aac	aac	cac	ata	atc	aca	aca
ser gly ile	att	lan	cer	ala	alıı	alv	leu	ile	leu	thr	asn	asn	his	val	ile	ala	ala
1021/341								1051	/351	_							
acc acc aad	cct	ccc	ctg	ggc	agt	ccg	ccg	ccg	aaa	acg	acg	gta	acc	ttc	tct	gac	aaa
ala ala lys	pro	pro	leu	gly	ser	pro	pro	pro	lys	thr	thr	val	thr	phe	ser	asp	gly
1081/361								1111			204	ast.	a + C	acc	ata	ata	cat
cgg acc gca arg thr ala	CCC	ttc	acg	gtg	gtg	ggg	gct	gac	nro	thr	ser	asp	ile	ala	val	val	arg
arg thr ala 1141/381	pro	pne	tnr	vaı	val	gry	ala	1171	./391	L	501	app		ala	• • • •		~~9
att caa aac	at.c	tcc	aaa	ctc	acc	ccq	atc	tcc	ctg	ggt	tcc	tcc	tcg	gac	ctg	agg	gtc
val gln gly	val	ser	gly	leu	thr	pro	ile	ser	leu	gly	ser	ser	ser	asp	leu	arg	val
1201/401								1233	./41:	L							
ggt cag ccg	gtg	ctg	gcg	atc	ggg	tcg	ccg	ctc	ggt	ttg	gag	ggc	acc	gtg	acc	acg	ggg
gly gln pro	val	leu	ala	ile	gly	ser	pro	Leu	gly 1/43	eu	gru	дтХ	thr	vai	Unr	CHI	gry
1261/421 atc gtc agc	aat	ctc	225	cat	cca	ata	tca				σασ	acc	aac	aac	caq	aac	acc
ile val ser	ala	leu	asn	arg	pro	val	ser	thr	thr	gly	glu	ala	gly	asn	gln	asn	thr
1321/441								135	L/45	1							
ata cta asc	gcc	att	cag	acc	gac	gcc	gcg	atc	aac	ccc	ggt	aac	tcc	ggg	ggc	gcg	ctg
val leu asp	ala	ile	gln	thr	asp	ala	ala	ile	asn	pro	gly	asn	ser	gly	gīā	ala	leu
1381/461									1/47		a++	acc	200	cta	aac	aca	aac
gtg aac atg	aac	gct	caa	CTC	gtc	gga	grc	aac	ser	ala	ile	ala	thr	leu	alv	ala	asp
1441/401								147	1/49	1							
tan acc ast	aca	caq	aqc	ggc	tcg	atc	ggt	ctc	ggt	ttt	gcg	att	cca	gtc	gac	cag	gcc
ser ala asp	ala	gln	ser	gly	ser	ile	gly	leu	gly	phe	ala	ile	pro	val	asp	gln	ala
1501/501								153	1/51	1							
aag cgc atc	gcc	gac	gag	ttg	atc	agc	acc	ggc	aag	gcg	tca	cat	gcc	tcc	ctg	ggt	grg
lys arg ile	ala	asp	glu	Leu	ıle	ser	thr	91y	1/53	а⊥а 1	ser	HITS	ата	261	160	gry	Vai
1561/521 cag gtg acc	+	~~~	222	aac	acc	cta	aac				ato	gaa	gta	ata	qcc	ggt	ggt
gln val thr	aau	. gac	lvs	asp	thr	leu	gly	ala	lys	ile	val	glu	val	val	āla	gly	gly
1621/5/1								165	1/55	1							
act acc acc	aac	gct	gga	gtg	ccg	aag	ggc	gto	gtt	gto	acc	aag	gto	gac	gac	cgc	ccg
ala ala ala	asn	ı ala	gly	val	pro	lys	gly	val	val	. val	. thr	lys	val	asp	asp	arç	pro
1681/561									1/57					י ממכ		acc	ata
atc aac ago	gcg	ggac	gcg	100	gtt	gec ala	gcc	g cg val	arc	r ser	· lvs	ala	pro	alı	, ala	thi	val
17/1/591								177	1/59	91							
aca cta aco	: ttt	cac	gat	ccc	: tcg	ggc	ggt	ago	: cgc	aca	gtg	caa	gto	acc	cto	ggd	aag
ala leu thi	r phe	glr	asp	pro	ser	gly	, gly	ser	arq	g thi	val:	gln	val	thi	: leu	ı gly	, lys
1801/601																	
gcg gag cag																	
ala glu glı	n OPA	4															

SEQ ID No.49F (continued 1)

FIGURE 49F (continued 1)

Fragment amplified by PCR based on the sequence similarities with a serine protease of the E.coli HtrA family (creation of an SnaBI site at the 3' end) and subcloned into the vector pJVEDa:

```
31/11
gat ccg gcg ggg cgg gtg tcg gcg cag gcg tgg ctg gcg gtc acg gcg gtg cgg gcg gtg
asp pro ala gly arg val ser ala gln ala trp leu ala val thr ala val arg ala val
                                      91/31
pro pro gly cys gly ala pro ala ala ala val ala met ala gly thr ala pro met pro
                                      151/51
121/41
aca tog toa gog gtg gag acg gtg goc tog gog gtg cog gtg gog gtg gog gat ggc tot
thr ser ser ala val glu thr val ala ser ala val pro val ala val ala asp gly ser
                                      211/71
181/61
thr ala thr ala gly pro ala asp thr ala asp lys ala gln ser ala ser ala ala ala
                                      271/91
241/81
ccq qcq gcg acg ggg gcc agg gcg gcg gcg gcc gcg gac tgt ggg gta ctg gcg gcg ccg
pro ala ala thr gly ala arg ala ala pro ala ala asp cys gly val leu ala ala pro
                                      331/111
301/101
geg gac acg geg ggc aag geg gtg gta eeg ggg gee eac ege tge eeg gte agg eag gea
ala asp thr ala gly lys ala val val pro gly ala his arg cys pro val arg gln ala
                                      391/131
361/121
tgg gcg ccg cgg gtg gcg ccg gtg ggc tga tcg gca acg gcg ggg ccg gcg gcg acg gcg
trp ala pro arg val ala pro val gly OPA ser ala thr ala gly pro ala ala thr ala
                                      451/151
421/141
gtg tcg gcg cgt ccg gcg ggg tcg ccg gag tag gcg gtg ccg gcg gga acg cca tgc tga
val ser ala arg pro ala gly ser pro glu AMB ala val pro ala gly thr pro cys OPA
                                      511/171
teg gge acg geg geg eeg geg eeg geg gag aca gea gtt teg eta atg geg egg eeg
ser gly thr ala ala pro ala ala pro ala glu thr ala val ser leu met ala arg pro
                                      571/191
541/181
gcg gcg cgg gcg gtg ccg gag ggc acc tct tcg gca atg gcg ggt ccg gcg gcc acg gcg
ala ala arg ala val pro glu gly thr ser ser ala met ala gly pro ala ala thr ala
601/201
                                      631/211
gag ccg tca cgg ccg gca aca ccg gta tcg gtg gcg ccg gcg gcg tcg gtg ggg acg cca
glu pro ser arg pro ala thr pro val ser val ala pro ala ala ser val gly thr pro
                                       691/231
661/221
ggc tga tcg gcc acg gtg gcg ccg gcg gtg ccg gcg ggg acc gcg ccg gag cct tgg ttg
gly OPA ser ala thr val ala pro ala val pro ala gly thr ala pro glu pro trp leu
                                       751/251
721/241
gcc gtg acg gcg ggc ccg gtg gga acg ggg gcg ctg gcg gcc agc tat acg gca acg gcg
ala val thr ala gly pro val gly thr gly ala leu ala ala ser tyr thr ala thr ala
                                      811/271
781/261
gcg acg gcg ccc ccg gca ccg gcg gaa cac tgc agg cgg cgg tga gcg gat tgg tga cgg
ala thr ala pro pro ala pro ala glu his cys arg arg OPA ala asp trp OPA arg
                                       871/291
841/281
ctt tgt tcg gtg cac ccg gcc aac ccg gcg aca ccg gcc aac ccg gct agc ccc gat caa
leu cys ser val his pro ala asn pro ala thr pro ala asn pro ala ser pro asp gln
                                      931/311
901/301
cga ggg ttt cgg tgc cgg tcc ggg gca tgg cca tcc gct gag ctg gcg atc tgg act acg
arg gly phe arg cys arg ser gly ala trp pro ser ala glu leu ala ile trp thr thr
                                       991/331
961/321
ttg gtg tag aaa aat cct gcc gcc cgg acc ctt aag gct ggg aca att tct gat agc tac
leu val AMB lys asn pro ala ala arg thr leu lys ala gly thr ile ser asp ser tyr
                                       1051/351
1021/341
ccc gac aca gga ggt tac ggg atg agc aat tcg cgc cgc cgc tca ctc agg tgg tca tgg
pro asp thr gly gly tyr gly met ser asn ser arg arg ser leu arg trp ser trp
                                       1111/371
1081/361
ttg ctg agc gtg ctg gct gcc gtc ggg ctg ggc ctg gcc acg gcg ccg gcc cag gcg gcc
leu leu ser val leu ala ala val gly leu gly leu ala thr ala pro ala gln ala ala
1141/381
ccg ccg gcc ttg tcg cag gac cgg tt
pro pro ala leu ser gln asp arg
```

SEQ ID No.50A

FIGURE 50A
REPLACEMENT SHEET (RILLE 26)



1 /1									31/11								
1/1 atc cgg	caa	aac	aaa	tat	caa	cac	agg	cat		caa	tca	caa	caa	tac	aaa	caa	tac
ile arg 61/21	arg	gly	gly	cys	arg	arg	arg	arg	gly trp 91/31	arg	ser	arg	arg	cys	gly	arg	cys
cgc cgg	gct	gtg	ggg	cgc	cgg	cgg	cgg	cgg	tgg caa	tgg	cgg	gaa	cgg	cgc	cga	tgc	caa
arg arg 121/41									151/51								
cat cgt his arg	cag	cgg	tgg	aga	cgg	tgg	cct	cgg	cgg tgc	cgg	tgg	cgg	trn	cgg	met	gct	leu
181/61	gin	arg	пр	ary	arg	CIP	pro	arg	211/71	arg	стр	419	CLP	urg	auc c	ulu	104
cgg cga	cgg	cgg	ggc	cgg	cgg	aca	cgg	cgg	aca agg	cgc	aat	cgg	cct	cgg	cgg	cgg	cgc
arg arg 241/81									271/91								
cgg cgg	cga	cgg	ggg	cca	āāā	cgg	cgc	cgg	ccg cgg	act	gtg	ggg	tac	tgg	cgg	cgc	cgg
arg arg 301/101	arg	arg	đтĀ	pro	дтА	arg	arg	arg	331/111		vai	gry	tyr	trp	arg	arg	arg
cgg aca	cgg	cgg	gca	agg	cgg	tgg	tac	cgg	ggg ccc	acc	gct	gcc	cgg	tca	ggc	agg	cat
arg thr 361/121	arg	arg	ala	arg	arg	trp	tyr	arg	gly pro 391/131		ala	ala	arg	ser	gly	arg	his
ggg cgc	cgc	ggg	tgg	cgc	cgg	tgg	gct	gat	cgg caa	cgg	cgg	ggc	cgg	cgg	cga	cgg	cgg
gly arg 421/141									451/151								
tgt cgg	cgc	gtc	cgg	cgg	ggt	cgc	cgg	agt	agg cgg	tgc	cgg	cgg	gaa	cgc	cat	gct	gat
cys arg 481/161									511/171								
cgg gca	cgg	cgg	cgc	cgg	cgg	cgc	cgg	cgg	aga cag	r cag	ttt	cgc	taa	tgg	cgc	aac	cgg
arg ala 541/181									571/191								
cgg <b>cg</b> c	ggg	cgg	tgc	cgg	agg	gca	cct	ctt	cgg caa	tgg	cgg	gtc	cgg	cgg	cca	cgg	cgg
arg arg 601/201									631/21:	-							
agc cgt	cac	ggc	cgg	caa	cac	cgg	tat	cgg	tgg cg	cgg	cgg	cgt	cgg	tgg	gga	cgc	cag
ser arg 661/221									691/23	L							
gct gat ala asp	cgg	cca	cgg	tgg	cgc	cgg	cgg	tgc	cgg cgg	g gga	ccg	cgc	cgg	agc	len	ggt	tgg
721/241									751/25	Ļ							
ccg tga	cgg	cgg	gcc	cgg	tgg	gaa	cgg	ggg	cgc tg	a caa	cca	gct	ata	cgg	caa	cgg	cgg
pro OPA 781/261									811/27	L							
cga cgg	cgc	ccc	cgg	cac	cgg	cgg	aac	act	gca gg	ggc	ggt	gag	cgg	att	ggt	gac	agc
arg arg 841/281									871/29	L							
ttt gtt	cgg	tgc	acc	cgg	cca	acc	cgg	cga	cac cg	g cca	acc	cgg	cta	gcc	ccg	atc	aac
phe val 901/301									931/31	L							
gag ggt	ttc	ggt	gcc	ggt	ccg	ggg	cat	ggc	cat cc	g ctc	ago	tgg	cga	tct	gga	cta	cgt
glu gly 961/321									991/33	1							
tgg tgt	aga	aaa	atc	ctg	ccg	ccc	gga	ccc	tta ag	g cto	gga	caa	ttt	ctg	ata	gct	acc
trp cys	1								1051/3	51							
ccg aca	cag	gag	gtt	acg	gga	tga	gca	att	cgc gc	c gcc	gct	cac	tca	ggt	ggt	cat	ggt
pro thr 1081/36	1								1111/3	71							
tgc tga	gcg	tgc	tgg	ctg	ccg	tcg	ggc	tgg	gcc tg	g cca	cgg	cgc	cgg	CCC	agg	cgg	pro
cys OPA 1141/38		cys	crp	теп	pro	ser	дтў	crp	ala tr	h hr	arc	, arg	ary	PLC	arg	arg	PLO
cgc cgg																	
arg arg	pro	cys	arg	arg	thr	gly	•										

SEQ ID No.50B

FIGURE 50B





31/11 tee gge ggg geg ggt gte gge gea gge gtg get gge ggt cae gge ggt geg ggt gee ser gly gly ala gly val gly ala gly val ala gly gly his gly gly ala gly gly ala 91/31 gcc ggg ctg tgg ggc gcc ggc ggc ggt ggc aat ggc ggg aac ggc gcc gat gcc aac ala gly leu trp gly ala gly gly gly gly asn gly gly asn gly ala asp ala asn 151/51 121/41 atc gtc agc ggt gga gac ggt ggc ctc ggc ggt gcc ggt ggc ggt ggc gga tgg ctc tac ile val ser gly gly asp gly gly leu gly gly ala gly gly gly gly gly trp leu tyr 211/71 181/61 ggc gac ggc ggg gcc ggc gga cac ggc gga caa ggc gca atc ggc ctc ggc ggc gcc gly asp gly gly ala gly gly his gly gly gln gly ala ile gly leu gly gly gly ala 271/91 241/81 ggc ggc gac ggg ggc cag ggc ggc gcc ggc cgc gga ctg tgg ggt act ggc ggc gcc ggc gly gly asp gly gly gln gly gly ala gly arg gly leu trp gly thr gly gly ala gly 331/111 gga cac ggc ggg caa ggc ggt ggt acc ggg ggc cca ccg ctg ccc ggt cag gca ggc atg gly his gly gly gln gly gly gly thr gly gly pro pro leu pro gly gln ala gly met 391/131 361/121 gly ala ala gly gly ala gly gly leu ile gly asn gly gly ala gly gly asp gly gly 451/151 gtc ggc gcg tcc ggc ggg gtc gcc gga gta ggc ggt gcc ggc ggg aac gcc atg ctg atc val gly ala ser gly gly val ala gly val gly gly ala gly gly asn ala met leu ile 511/171 481/161 gly his gly gly ala gly gly ala gly gly asp ser ser phe ala asn gly ala ala gly 571/191 541/181 ggc gcg ggc ggt gcc gga ggg cac ctc ttc ggc aat ggc ggg tcc ggc ggc cac ggc gga gly ala gly gly ala gly gly his leu phe gly asn gly gly ser gly gly his gly gly 631/211 gcc gtc acg gcc ggc aac acc ggt atc ggt ggc gcc ggc ggc gtc ggt ggg gac gcc agg ala val thr ala gly asn thr gly ile gly gly ala gly gly val gly gly asp ala arg 691/231 661/221 ctg atc ggc cac ggt ggc gcc ggc ggt gcc ggc ggg gac cgc gcc gga gcc ttg gtt ggc leu ile gly his gly gly ala gly gly ala gly gly asp arg ala gly ala leu val gly 751/251 721/241 cgt gac ggc ggg ccc ggt ggg aac ggg ggc gct ggc ggc cag cta tac ggc aac ggc ggc arg asp gly gly pro gly gly asn gly gly ala gly gly gln leu tyr gly asn gly gly 811/271 781/261 gac ggc gcc ccc ggc acc ggc gga aca ctg cag gcg gcg gtg agc gga ttg gtg acg gct asp gly ala pro gly thr gly gly thr leu gln ala ala val ser gly leu val thr ala 871/291 841/281 ttg ttc ggt gca ccc ggc caa ccc ggc gac acc ggc caa ccc ggc tag ccc cga tca acg leu phe gly ala pro gly gln pro gly asp thr gly gln pro gly AMB pro arg ser thr 931/311 901/301 agg gtt tog gtg cog gto ogg ggo atg goo ato ogo tga got ggo gat otg gao tao gtt arg val ser val pro val arg gly met ala ile arg OPA ala gly asp leu asp tyr val 991/331 961/321 ggt gta gaa aaa tcc tgc cgc ccg gac cct taa ggc tgg gac aat ttc tga tag cta ccc gly val glu lys ser cys arg pro asp pro OCH gly trp asp asn phe OPA AMB leu pro 1051/351 1021/341 cga cac agg agg tta cgg gat gag caa ttc gcg ccg ccg ctc act cag gtg gtc atg gtt arg his arg arg leu arg asp glu gln phe ala pro pro leu thr gln val val met val 1111/371 1081/361 get gag egt get gge tge egt egg get ggg eet gge eac gge gee gge eea gge eec ala glu arg ala gly cys arg arg ala gly pro gly his gly ala gly pro gly pro 1141/381 gcc ggc ctt gtc gca gga ccg gtt ala gly leu val ala gly pro val

SEQ ID No.50C

FIGURE 50C





Coding sequence Rv0125 predicted by Cole et al., 1998 (Nature 393:537-544) and containing seq50A:

1/1									31/1	1								
ato aoc	aat	tcg	cqc	cgc	cgc	tca	ctc	agg	tgg	tca	tgg	ttg	ctg	agc	gtg	ctg	gct	gcc
Met ser 61/21	asn	ser	arg	arg	arg	ser	leu	arg	trp 91/3	ser	trp	leu	leu	ser	val	leu	ala	ala
atc aga	cta	ggc	ctg	gcc	acg	gcg	ccg	gcc	cag	gcg	gcc	ccg	ccg	gcc	ttg	tcg	cag	gac
val gly	leu	gly	leu	ala	thr	ala	pro	ala	gln 151/	ala 51	ala	pro	pro	ala	leu	ser	gln	asp
caa ttc	gcc	gac	ttc	ccc	gcg	ctg	ccc	ctc	gac	ccg	tcc	gcg	atg	gtc	gcc	caa	gtg	ggg
arg phe 181/61									211/	71								
cca cag	gtg	gtc	aac	atc	aac	acc	aaa	ctg	ggc	tac	aac	aac	gcc	gtg	ggc	gcc	āāā	acc
pro gln 241/81									271/	91								
ggc atc	gtc	atc	gat	CCC	aac	ggt	gtc	gtg	ctg	acc	aac	aac	cac	gtg	atc	gcg	ggc	gcc
gly ile 301/101									331/	111								
acc gac	atc	aat	gcg	ttc	agc	gtc	ggc	tcc	ggc	caa	acc	tac	ggc	gtc	gat	gtg	gtc	ggg
thr asp 361/121									391/	131								
tat gac	cgc	acc	cag	gat	gtc	aca	gtg	ctg	cag	ctg	cgc	ggt	gcc	ggt	ggc	ctg	ccg	tcg
tyr asp 421/141									451/	151								
gcg gcg	atc	ggt	ggc	ggc	gtc	gcg	gtt	ggt	gag	ccc	gtc	gtc	gcg	atg	ggc	aac	agc	ggt
ala ala 481/161									511/	171								
ggg cag	ggc	gga	acg	ccc	cgt	gcg	gtg	cct	ggc	agg	gtg	gtc	gcg	ctc	ggc	caa	acc	gtg
gly gln 541/181									571/	191								
cag gcg	tcg	gat	tcg	ctg	acc	ggt	gcc	gaa	gag	aca	ttg	aac	ggg	ttg	atc	cag	ttc	gat
gln ala 601/201									631/	/211								
gcc gcg	atc	cag	CCC	ggt	gat	tcg	ggc	ggg	CCC	gtc	gtc	aac	ggc	cta	gga	cag	gtg	gtc
ala ala 661/221									691/	/231								
ggt atg	aac	acg	gcc	gcg	tcc	gat	aac	ttc	cag	Lou	CON	cag aln	ggu	999	aln	gya gya	nhe	ala
721/241									751,	/251								
att ccg	atc	ggg	cag	gcg	atg	gcg	atc	gcg	ggc	cag	ile	ara	ser	ggu	ggg	gyy	ser	pro
781/261		gry	gin	ата	mec	ата	110	ara		/271		urg	501	9-1	9-1	5-1		F
acc gtt	cat	ato	aaa	cct	acc	gcc	ttc	ctc	ggc	ttg	ggt	gtt	gtc	gac	aac	aac	ggc	aac
thr val	his	ile	gly	pro	thr	ala	phe	leu	gly	leu	gly	val	val	asp	asn	asn	gly	asn
841/281									871.	/291								
ggc gca	ı cga	gto	caa	. cgc	gtg	gto	ggg	ago	gct	ccg	gcg	gca	agt	ctc	ggc	ato	COL	thr
gly ala 901/301									931	/311								
ggc gac	gtg	ato	acc	gcg	gto	gac	ggc	gct	ccg	ato	aac	tcg	gcc	acc	gcg	atg	gcc	gac
gly asp		. ile	thr	ala	. vaı	asp	g T	ala	991	11e 331/	ası	ı ser	ala	. CIII	ата	i ille c	arc	asp
961/321 gcg ctt	aac	aac	cat	cat	ccc	: aat	gac	gto	atc	tcg	gto	acc	tgg:	caa	acc	aag	tcg	ggc
ala le	ı asr	alv	his	his	pro	gly	asp	val	ile	ser	: val	thr	trp	glr	thr	lys	sei	gly
1021/34	11								105	1/35	1							
ggc ac	g cgt	aca	ggg	g aac	gtg	aca	ttg	l dcc	gag	gga	CCC	ccg	gco	tga	l			
gly th	rarç	g thr	gly	asr	ı val	. thi	leu	ala	glu	ğΤ?	pro	pro	ата	OPA	1			

SEQ ID No.50D

FIGURE 50D

OPA





#### 180/185

ORF according to Cole et al., 1998 (Nature 393:537-544) and containing Rv0125:

```
31/11
1/1
tag aaa aat oot goo goo ogg acc ott aag got ggg aca att tot gat ago tac ooc gac
AMB lys asn pro ala ala arg thr leu lys ala gly thr ile ser asp ser tyr pro asp
                                        91/31
61/21
aca gga ggt tac ggg atg agc aat tcg cgc cgc cgc tca ctc agg tgg tca tgg ttg ctg
thr gly gly tyr gly met ser asn ser arg arg ser leu arg trp ser trp leu leu
                                        151/51
121/41
age gtg ctg gct gcc gtc ggg ctg ggc ctg gcc acg gcg ccg gcc cag gcg ccg ccg
ser val leu ala ala val gly leu gly leu ala thr ala pro ala gln ala ala pro pro
                                        211/71
181/61
gcc ttg tcg cag gac cgg ttc gcc gac ttc ccc gcg ctg ccc ctc gac ccg tcc gcg atg
ala leu ser gln asp arg phe ala asp phe pro ala leu pro leu asp pro ser ala met
                                        271/91
gtc gcc caa gtg ggg cca cag gtg gtc aac atc aac acc aaa ctg ggc tac aac aac gcc
val ala gin val gly pro gln val val asn ile asn thr lys leu gly tyr asn asn ala
                                        331/111
gtg ggc gcc ggg acc ggc atc gtc atc gat ccc aac ggt gtc gtg ctg acc aac aac cac
val gly ala gly thr gly ile val ile asp pro asn gly val val leu thr asn asn his
                                        391/131
gtg atc gcg ggc gcc acc gac atc aat gcg ttc agc gtc ggc tcc ggc caa acc tac ggc
val ile ala gly ala thr asp ile asn ala phe ser val gly ser gly gln thr tyr gly
                                        451/151
gtc gat gtg gtc ggg tat gac cgc acc cag gat gtc gcg gtg ctg cag ctg cgc ggt gcc
val asp val val gly tyr asp arg thr gln asp val ala val leu gln leu arg gly ala
                                        511/171
ggt ggc ctg ccg tcg gcg gcg atc ggt ggc gtc gcg gtt ggt gag ccc gtc gtc gcg
gly gly leu pro ser ala ala ile gly gly val ala val gly glu pro val val ala
                                        571/191
541/181
atg ggc aac agc ggt ggg cag ggc gga acg ccc cgt gcg gtg cct ggc agg gtg gtc gcg
met gly asn ser gly gly gln gly gly thr pro arg ala val pro gly arg val val ala
                                         631/211
601/201
ctc ggc caa acc gtg cag gcg tcg gat tcg ctg acc ggt gcc gaa gag aca ttg aac ggg
leu gly gln thr val gln ala ser asp ser leu thr gly ala glu glu thr leu asn gly
                                         691/231
661/221
ttg atc cag ttc gat gcc gcg atc cag ccc ggt gat tcg ggc ggg ccc gtc gtc aac ggc
leu ile gln phe asp ala ala ile gln pro gly asp ser gly gly pro val val asn gly
                                         751/251
721/241
cta gga cag gtg gtc ggt atg aac acg gcc gcg tcc gat aac ttc cag ctg tcc cag ggt
leu gly gln val val gly met asn thr ala ala ser asp asn phe gln leu ser gln gly
                                         811/271
781/261
ggg cag gga ttc gcc att ccg atc ggg cag gcg atg gcg atc gcg ggc cag atc cga tcg
 gly gln gly phe ala ile pro ile gly gln ala met ala ile ala gly gln ile arg ser
                                         871/291
 841/281
 ggt ggg ggg tca ccc acc gtt cat atc ggg cct acc gcc ttc ctc ggc ttg ggt gtt gtc
gly gly gly ser pro thr val his ile gly pro thr ala phe leu gly leu gly val val
                                         931/311
 901/301
 gac aac aac ggc aac ggc gca cga gtc caa cgc gtg gtc ggg agc gct ccg gcg gca agt
 asp asn asn gly asn gly ala arg val gln arg val val gly ser ala pro ala ala ser
                                         991/331
 961/321
 ctc ggc atc tcc acc ggc gac gtg atc acc gcg gtc gac ggc gct ccg atc aac tcg gcc
 leu gly ile ser thr gly asp val ile thr ala val asp gly ala pro ile asn ser ala
                                         1051/351
 acc gcg atg gcg gac gcg ctt aac ggg cat cat ccc ggt gac gtc atc tcg gtg acc tgg
 thr ala met ala asp ala leu asn gly his his pro gly asp val ile ser val thr trp
                                         1111/371
 1081/361
 caa acc aag tog ggc ggc acg cgt aca ggg aac gtg aca ttg gcc gag gga ccc ccg gcc
 gln thr lys ser gly gly thr arg thr gly asn val thr leu ala glu gly pro pro ala
 1141/381
 tga
```

SEQ ID No.50F

FIGURE 50F



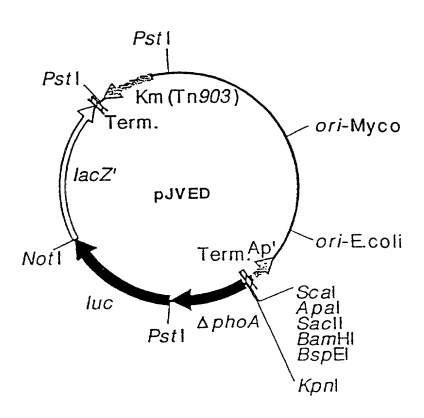


FIGURE 51A

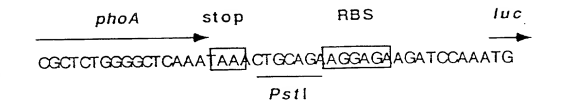
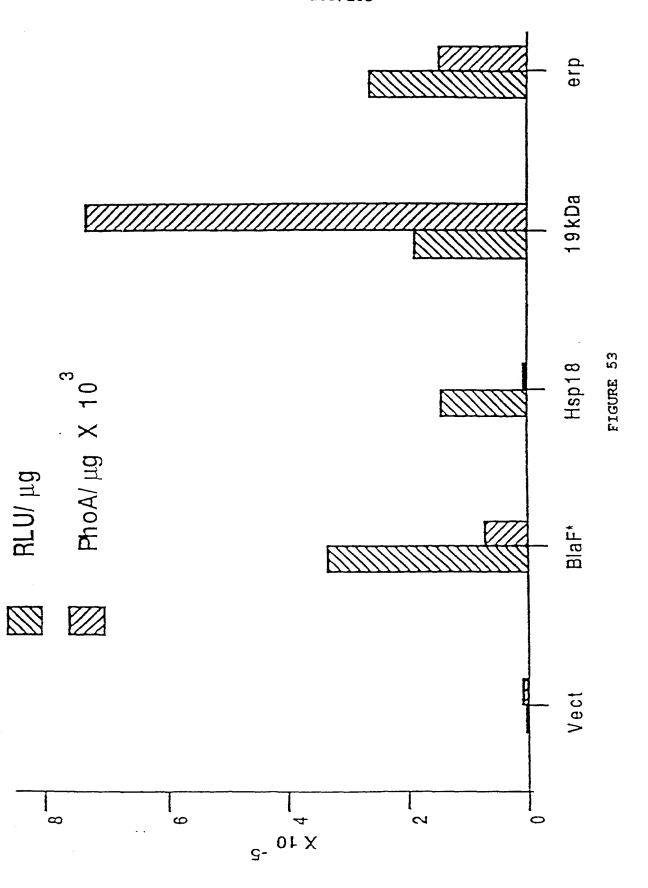


FIGURE 51B



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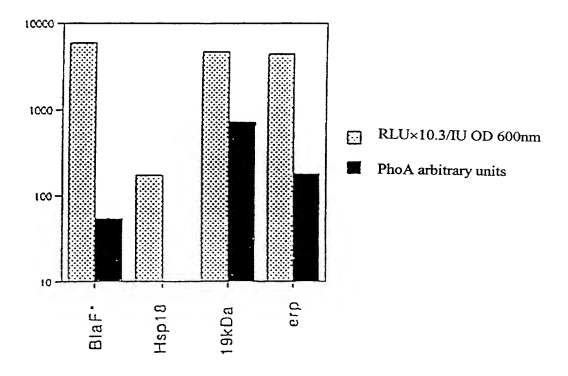
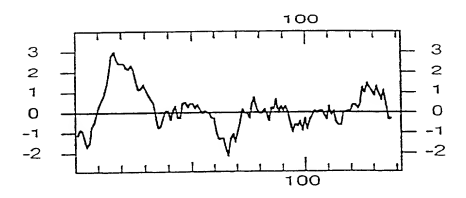


FIGURE 54



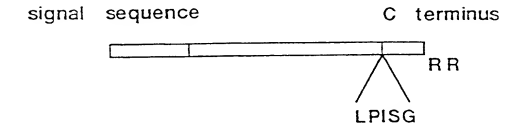


FIGURE 55
REPLACEMENT SHEET (RULE 26)

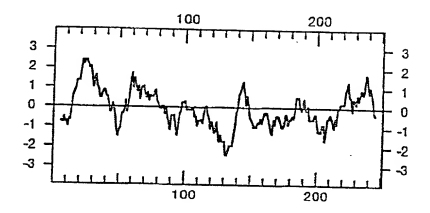


FIGURE 56

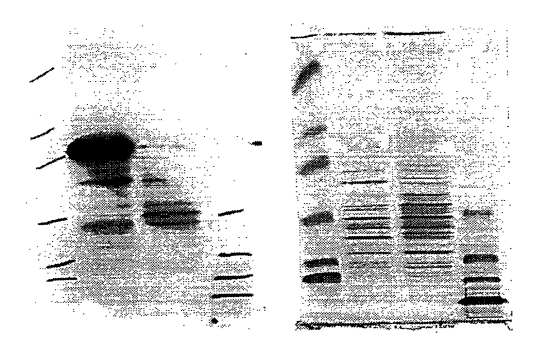


FIGURE 57A

FIGURE 57B